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94043 (US). RANK, David, R. [US/US]; 117 El Dorado Commons, Fremont, CA 94539 (US).

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(74) Agent: RONNING, Royal, N., Jr.; Amersham Pharma-cia Biotech, Inc., 800 Centennial Avenue, Piscataway, NJ 08855 (US).

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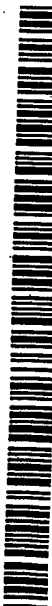
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**WO 01/57275 A2**

(54) Title: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FOR ANALYSIS OF GENE EXPRESSION IN HUMAN BRAIN

(57) Abstract: A single exon nucleic acid microarray comprising a plurality of single exon nucleic acid probes for measuring gene expression in a sample derived from human brain is described. Also described are single exon nucleic acid probes expressed in the brain and their use in methods for detecting gene expression.

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HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL  
FOR ANALYSIS OF GENE EXPRESSION IN HUMAN BRAIN

CROSS REFERENCE TO RELATED APPLICATIONS

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The present application is a continuation-in-part of U.S. patent application serial nos. 09/632,366, filed August 3, 2000 and 09/608,408, filed June 30, 2000; claims the benefit under 35 U.S.C. s 119(e) of U.S. provisional patent 10 application serial nos. 60/236,359, filed September 27, 2000, 60/234,687, filed September 21, 2000, 60/207,456, filed May 26, 2000, and 60/180,312, filed February 4, 2000; and further claims the benefit under 35 U.S.C. s 119(a) of UK patent application no. 0024263.6, filed October 4, 2000, 15 the disclosures of which are incorporated herein by reference in their entireties.

REFERENCE TO SEQUENCE LISTING AND INCORPORATION BY  
REFERENCE THEREOF

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The present application includes a Sequence Listing in electronic format, filed pursuant to PCT Administrative Instructions 801 - 806 on a single CD-R disc, in triplicate, containing a file named pto\_BRAIN.txt, created 25 24 January 2001, having 25,840,972 bytes. The Sequence Listing contained in said file on said disc is incorporated herein by reference in its entirety.

Field of the Invention

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The present invention relates to genome-derived single exon microarrays useful for verifying the expression of regions of genomic DNA predicted to encode protein. In particular, the present invention relates to unique genome-derived single exon nucleic acid probes expressed in human

brain and single exon nucleic acid microarrays that include such probes.

Background of the Invention

5       For almost two decades following the invention of general techniques for nucleic acid sequencing, Sanger *et al.*, *Proc. Natl. Acad. Sci. USA* 70(4):1209-13 (1973); Gilbert *et al.*, *Proc. Natl. Acad. Sci. USA* 70(12):3581-4 (1973), these techniques were used principally as tools to  
10 further the understanding of proteins – known or suspected – about which a basic foundation of biological knowledge had already been built. In many cases, the cloning effort that preceded sequence identification had been both informed and directed by that antecedent  
15 biological understanding.

For example, the cloning of the T cell receptor for antigen was predicated upon its known or suspected cell type-specific expression, by its suspected membrane association, and by the predicted assembly of its gene via  
20 T cell-specific somatic recombination. Subsequent sequencing efforts at once confirmed and extended understanding of this family of proteins. Hedrick *et al.*, *Nature* 308(5955):153-8 (1984).

More recently, however, the development of high  
25 throughput sequencing methods and devices, in concert with large public and private undertakings to sequence the human and other genomes, has altered this investigational paradigm: today, sequence information often precedes understanding of the basic biology of the encoded protein  
30 product.

One of the approaches to large-scale sequencing is predicated upon the proposition that expressed sequences – that is, those accessible through isolation of mRNA – are of greatest initial interest. This "expressed  
35 sequence tag" ("EST") approach has already yielded vast

amounts of sequence data (see for example Adams *et al.*, *Science* 252:1651 (1991); Williamson, *Drug Discov. Today* 4:115 (1999)). For nucleic acids sequenced by this approach, often the only biological information that is  
5 known *a priori* with any certainty is the likelihood of biologic expression itself. By virtue of the species and tissue from which the mRNA had originally been obtained, most such sequences are also annotated with the identity of the species and at least one tissue in which expression  
10 appears likely.

More recently, the pace of genomic sequencing has accelerated dramatically. When genomic DNA serves as the initial substrate for sequencing efforts, expression cannot be presumed; often the only *a priori* biological information  
15 about the sequence includes the species and chromosome (and perhaps chromosomal map location) of origin.

With the ever-accelerating pace of sequence accumulation by directed, EST, and genomic sequencing approaches – and in particular, with the accumulation of  
20 sequence information from multiple genera, from multiple species within genera, and from multiple individuals within a species – there is an increasing need for methods that rapidly and effectively permit the functions of nucleic sequences to be elucidated. And as such functional  
25 information accumulates, there is a further need for methods of storing such functional information in meaningful and useful relationship to the sequence itself; that is, there is an increasing need for means and apparatus for annotating raw sequence data with known or  
30 predicted functional information.

Although the increase in the pace of genomic sequencing is due in large part to technological changes in sequencing strategies and instrumentation, Service, *Science* 280:995 (1998); Pennisi, *Science* 283: 1822-1823 (1999),  
35 there is an important functional motivation as well.

While it was understood that the EST approach would rarely be able to yield sequence information about the noncoding portions of the genome, it now also appears the EST approach is capable of capturing only a fraction of 5 a genome's actual expression complexity.

For example, when the *C. elegans* genome was fully sequenced, gene prediction algorithms identified over 19,000 potential genes, of which only 7,000 had been found by EST sequencing. *C. elegans* Sequencing Consortium, 10 *Science* 282:2012 (1998). Analogously, the recently completed sequence of chromosome 2 of *Arabidopsis* predicts over 4000 genes, Lin et al., *Nature*, 402:761 (1999), of which only about 6% had previously been identified via EST sequencing efforts. Although the human genome has the 15 greatest depth of EST coverage, it is still woefully short of surrendering all of its genes. One recent estimate suggests that the human genome contains more than 146,000 genes, which would at this point leave greater than half of the genes undiscovered. It is now predicted that many 20 genes, perhaps 20 to 50%, will only be found by genomic sequencing.

There is, therefore, a need for methods that permit the functional regions of genomic sequence – and most importantly, but not exclusively, regions that 25 function to encode genes – to be identified.

Much of the coding sequence of the human genome is not homologous to known genes, making detection of open reading frames ("ORFs") and predictions of gene function difficult. Computational methods exist for predicting 30 coding regions in eukaryotic genomes. Gene prediction programs such as GRAIL and GRAIL II, Uberbacher et al., *Proc. Natl. Acad. Sci. USA* 88(24):11261-5 (1991); Xu et al., *Genet. Eng.* 16:241-53 (1994); Uberbacher et al., *Methods Enzymol.* 266:259-81 (1996); GENEFINDER, Solovyev et 35 al., *Nucl. Acids. Res.* 22:5156-63 (1994); Solovyev et al.,

Ismb 5:294-302 (1997); and GENESCAN, Burge et al., *J. Mol. Biol.* 268:78-94 (1997), predict many putative genes without known homology or function. Such programs are known, however, to give high false positive rates. Burset et al., 5 *Genomics* 34:353-367 (1996). Using a consensus obtained by a plurality of such programs is known to increase the reliability of calling exons from genomic sequence. Ansari-Lari et al., *Genome Res.* 8(1):29-40 (1998)

Identification of functional genes from genomic 10 data remains, however, an imperfect art. For example, in reporting the full sequence of human chromosome 21, the Chromosome 21 Mapping and Sequencing Consortium reports that prior bioinformatic estimates of human gene number may need to be revised substantially downwards. *Nature* 15 405:311-199 (2000); Reeves, *Nature* 405:283-284 (2000).

Thus, there is a need for methods and apparatus that permit the functions of the regions identified bioinformatically - and specifically, that permit the expression of regions predicted to encode protein - readily 20 to be confirmed experimentally.

Recently, the development of nucleic acid microarrays has made possible the automated and highly parallel measurement of gene expression. Reviewed in Schena (ed.), DNA Microarrays : A Practical Approach 25 (Practical Approach Series), Oxford University Press (1999) (ISBN: 0199637768); *Nature Genet.* 21(1) (suppl):1 - 60 (1999); Schena (ed.), Microarray Biochip: Tools and Technology, Eaton Publishing Company/BioTechniques Books Division (2000) (ISBN: 1881299376).

It is common for microarrays to be derived from cDNA/EST libraries, either from those previously described in the literature, such as those from the I.M.A.G.E. consortium, Lennon et al., *Genomics* 33(1):151-2 (1996), or from the construction of "problem specific" libraries 30 35 targeted at a particular biological question, R.S. Thomas

et al., *Cancer Res.* (in press). Such microarrays by definition can measure expression only of those genes found in EST libraries, and thus have not been useful as probes for genes discovered solely by genomic sequencing.

- 5       The utility of using whole genome nucleic acid microarrays to answer certain biological questions has been demonstrated for the yeast *Saccharomyces cerevisiae*. De Risi et al., *Science* 278:680 (1997). The vast majority of yeast nuclear genes, approximately 95% however, are single 10 exon genes, i.e., lack introns, Lopez et al., *RNA* 5:1135-1137 (1999); Goffeau et al., *Science* 274:563-67 (1996), permitting coding regions more readily to be identified. Whole genome nucleic acid microarrays have not generally been used to probe gene expression from more complex 15 eukaryotic genomes, and in particular from those averaging more than one intron per gene.

- Diseases of the brain and nervous system are a significant cause of human morbidity and mortality. Increasingly, genetic factors are being found that 20 contribute to predisposition, onset, and/or aggressiveness of most, if not all, of these diseases. Although mutations in single genes have been identified as causative for some diseases of the brain and nervous system, for the most part these disorders are believed to have polygenic etiologies. 25 There is a need for methods and apparatus that permit prediction, diagnosis and prognosis of diseases of the brain and nervous system particularly those diseases with polygenic etiology.

30 Summary of the Invention

The present invention solves these and other problems in the art by providing methods and apparatus for predicting, confirming, and displaying functional 35 information derived from genomic sequence. The present

invention also provides apparatus for verifying the expression of putative genes identified within genomic sequence.

In particular, the invention provides novel 5 genome-derived single exon nucleic acid microarrays useful for verifying the expression of putative genes identified within genomic sequence.

The present invention also provides compositions and kits for the ready production of nucleic acids 10 identical in sequence to, or substantially identical in sequence to, probes on the genome-derived single exon microarrays of the present invention.

Accordingly, in a first aspect of the invention, there is provided a spatially-addressable set of single 15 exon nucleic acid probes for measuring gene expression in a sample derived from human brain, comprising a plurality of single exon nucleic acid probes according to any one of the nucleotide sequences set out in SEQ ID NOS: 1 - 12,821 or a complementary sequence, or a portion of such a sequence.

20 By plurality is meant at least two, suitably at least 20, most suitably at least 100, preferably at least 1000 and, most preferably, upto 5000.

In one embodiment of the first aspect, each of said plurality of probes is separately and addressably 25 amplifiable.

In an alternative embodiment, each of said plurality of probes is separately and addressably isolatable from said plurality.

30 In a preferred embodiment, each of said plurality of probes is amplifiable using at least one common primer. Preferably, each of said plurality of probes is amplifiable using a first and a second common primer.

In yet another embodiment, said set of single exon nucleic acid probes comprises between 50 - 20,000 35 probes, for example, 50 - 5000.

Suitably, said set of single exon nucleic acid probes comprises at least 50 - 1000 discrete single exon nucleic acid probes having a sequence as set out in any of SEQ ID NOS.: 1 - 25,434 or a complimentary sequence, or a portion of such a sequence.

Preferably, the average length of the single exon nucleic acid probes is between 200 and 500 bp. It is preferred that the average length should be at least 200bp, suitably at least 250bp, most suitably at least 300bp, preferably at least 400bp and, most preferably, 500 bp.

In another embodiment, the single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence. It is preferred that at least 50%, suitably at least 60%, most suitably at least 70%, preferably at least 75%, more preferably at least 80, 85, 90, 95 or 99% of said single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence.

In another preferred embodiment, said single exon nucleic acid lack homopolymeric stretches of A or T. It is preferred that at least 50%, suitably at least 60%, most suitably at least 70%, preferably at least 75%, more preferably at least 80, 85, 90, 95 or 99% of said single exon nucleic acid probes lack homopolymeric stretches of A or T.

Preferably, a spatially-addressable set of single exon nucleic acid probes in accordance with the first aspect of the invention is addressably disposed upon a substrate.

Suitable substrates include a filter membrane which may, preferably, be nitrocellulose or nylon. The nylon may preferably, be positively-charged. Other suitable substrates include glass, amorphous silicon, crystalline silicon, and plastic. Further suitable materials include polymethylacrylic, polyethylene, polypropylene, polyacrylate, polymethylmethacrylate, polyvinylchloride,

polytetrafluoroethylene, polystyrene, polycarbonate, polyacetal, polysulfone, celluloseacetate, cellulosenitrate, nitrocellulose, and mixtures thereof.

In a second aspect of the invention, there is  
5 provided a microarray comprising a spatially addressable set of single exon nucleic acid probes in accordance with the first aspect of the invention.

In one embodiment, a genome-derived single-exon microarray is packaged together with such an ordered set of 10 amplifiable probes corresponding to the probes, or one or more subsets of probes, thereon. In alternative embodiments, the ordered set of amplifiable probes is packaged separately from the genome-derived single exon microarray.

15 In another aspect, the invention provides genome-derived single exon nucleic acid probes useful for gene expression analysis, and particularly for gene expression analysis by microarray. In particular embodiments of this aspect, the present invention provides human single-exon 20 probes that include specifically-hybridizable fragments of SEQ ID Nos. 12,822 - 25,434, wherein the fragment hybridizes at high stringency to an expressed human gene. In particular embodiments, the invention provides single exon probes comprising SEQ ID Nos. 1 - 12,821.

25 Accordingly, in a third aspect of the invention, there is provided a single exon nucleic acid probe for measuring human gene expression in a sample derived from human brain which is a nucleic acid molecule comprising a nucleotide sequence as set out in any of SEQ ID NOS.: 1 - 30 12,821 or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid expressed in the human brain.

In one embodiment, a single exon nucleic acid probe in accordance with the third aspect comprises a 35 nucleotide sequence as set out in any of SEQ ID NOS.:

12,822 - 25,434 or a complementary sequence or a fragment thereof.

In a fourth aspect of the invention, there is provided a single exon nucleic acid probe for measuring 5 human gene expression in a sample derived from human brain which is a nucleic acid molecule having a sequence encoding a peptide comprising a peptide sequence as set out in any of SEQ ID NOS.: 25,435 - 37,811 or a complementary sequence or a fragment thereof wherein said probe hybridizes at high 10 stringency to a nucleic acid expressed in the human brain.

Preferably, a single exon nucleic acid probe in accordance with the third or fourth aspects of the invention comprises between at least 15 and 50 contiguous nucleotides of said SEQ ID NO:. It is preferred that the 15 single exon nucleic acid probe comprises at least 15, suitably at least 20, more suitably at least 25 or preferably at least 50 contiguous nucleotides of said SEQ ID NO:.

In another preferred embodiment, a single exon 20 nucleic acid probe in accordance with the third or fourth aspects of the invention is between 3kb and 25kb in length. It is preferred that said probe is no more than 3kb, suitably no more than 5kb, more suitably no more than 10kb, preferably 15kb, more preferably 20kb or, most preferably, 25 no more than 20kb in length.

Preferably, a single exon nucleic acid probe in accordance with either the fifth or sixth aspect of the invention is DNA, preferably single-stranded DNA, RNA or PNA.

30 In another embodiment of either the third or fourth aspect of the invention, a single exon nucleic acid probe is detectably labeled. Suitable detectable labels include a radionuclide, a fluorescent label or a first member of a specific binding pair. Suitable fluorescent 35 labels include dyes such as cyanine dyes, preferably Cy3

and Cy5 although other suitable dyes will be known to those skilled in the art.

In a particularly preferred embodiment, a single exon nucleic acid probe in accordance with either the third or fourth aspect of the invention lacks prokaryotic and bacteriophage vector sequence. In yet another embodiment, a single exon nucleic acid probe in accordance with either the third or fourth aspect of the invention lacks homopolymeric stretches of A or T.

10 In a fifth aspect of the invention, there is provided an amplifiable nucleic acid composition, comprising:

the single exon nucleic acid probe in accordance with either of the third or fourth aspects of the  
15 invention; and at least one nucleic acid primer;

wherein said at least one primer is sufficient to prime enzymatic amplification of said probe.

20 In an sixth aspect of the invention, there is provided a method of measuring gene expression in a sample derived from human brain, comprising:

25 contacting the single exon microarray in accordance with the second aspect of the invention, with a first collection of detectably labeled nucleic acids, said first collection of nucleic acids derived from mRNA of human brain; and then

measuring the label detectably bound to each probe of said microarray.

30 In a seventh aspect of the invention, there is provided a method of identifying exons in a eukaryotic genome, comprising:

algorithmically predicting at least one exon from genomic sequence of said eukaryote; and then

detecting specific hybridization of detectably labeled nucleic acids to a single exon probe,

35 wherein said detectably labeled nucleic acids are

derived from mRNA from the brain of said eukaryote, said probe is a single exon probe having a fragment identical in sequence to, or complementary in sequence to, said predicted exon, said probe is included within a single exon 5 microarray in accordance with the first aspect of the invention, and said fragment is selectively hybridizable at high stringency..

In a eighth aspect of the invention, there is provided a method of assigning exons to a single gene, 10 comprising:

identifying a plurality of exons from genomic sequence in accordance with the seventh aspect of the invention; and then

measuring the expression of each of said exons in 15 a plurality of tissues and/or cell types using hybridization to single exon microarrays having a probe with said exon,

wherein a common pattern of expression of said exons in said plurality of tissues and/or cell types 20 indicates that the exons should be assigned to a single gene.

In an ninth aspect of the invention, there is provided a nucleic acid sequence as set out in any of SEQ ID NOS: 1 - 25,434 wherein said sequence encodes a peptide. 25

In a tenth aspect of the invention, there is provided a peptide encoded by a sequence comprising a sequence as set out in any of SEQ ID NOS: 12,822 - 25,434, or a complementary sequence or coding portion thereof.

In a preferred embodiment, a peptide may be 30 encoded by a sequence comprising a sequence set out in any of SEQ ID NOS.: 1 - 12,821.

In a further aspect, the invention provides peptides comprising an amino acid sequence translated from the DNA fragments, said amino acid sequences comprising SEQ 35 ID NOS.: 25,435 - 37,811.

Accordingly in a eleventh aspect of the invention there is provided a peptide comprising a sequence as set out in any of SEQ ID NOS: 25,435 - 37,811, or fragment thereof.

- 5        In another aspect, the invention provides means for displaying annotated sequence, and in particular, for displaying sequence annotated according to the methods and apparatus of the present invention. Further, such display can be used as a preferred graphical user interface for  
10      electronic search, query, and analysis of such annotated sequence.

#### Detailed Description of the Invention

15

##### Definitions

As used herein, the term "microarray" and phrase "nucleic acid microarray" refer to a substrate-bound collection of plural nucleic acids, hybridization to each 20 of the plurality of bound nucleic acids being separately detectable. The substrate can be solid or porous, planar or non-planar, unitary or distributed.

As so defined, the term "microarray" and phrase "nucleic acid microarray" include all the devices so called 25 in Schena (ed.), DNA Microarrays: A Practical Approach (Practical Approach Series), Oxford University Press (1999) (ISBN: 0199637768); *Nature Genet.* 21(1) (suppl):1 - 60 (1999); and Schena (ed.), Microarray Biochip: Tools and Technology, Eaton Publishing Company/BioTechniques Books 30 Division (2000) (ISBN: 1881299376). As so defined, the term "microarray" and phrase "nucleic acid microarray" further include substrate-bound collections of plural 35 nucleic acids in which the nucleic acids are distributably disposed on a plurality of beads, rather than on a unitary planar substrate, as is described, *inter alia*, in Brenner

et al., Proc. Natl. Acad. Sci. USA 97(4):166501670 (2000); in such case, the term "microarray" and phrase "nucleic acid microarray" refer to the plurality of beads in aggregate.

- 5 As used herein with respect to a nucleic acid microarray, the term "probe" refers to the nucleic acid that is, or is intended to be, bound to the substrate; in such context, the term "target" thus refers to nucleic acid intended to be bound thereto by Watson-Crick  
10 complementarity. As used herein with respect to solution phase hybridization, the term "probe" refers to the nucleic acid of known sequence that is detectably labeled.

- As used herein, the expression "probe comprising SEQ ID NO.", and variants thereof, intends a nucleic acid  
15 probe, at least a portion of which probe has either (i) the sequence directly as given in the referenced SEQ ID NO., or (ii) a sequence complementary to the sequence as given in the referenced SEQ ID NO., the choice as between sequence directly as given and complement thereof dictated by the  
20 requirement that the probe hybridize to mRNA.

- As used herein, the term "open reading frame" and the equivalent acronym "ORF" refer to that portion of an exon that can be translated in its entirety into a sequence of contiguous amino acids i.e. a nucleic acid sequence  
25 that, in at least one reading frame, does not possess stop codons; the term does not require that the ORF encode the entirety of a natural protein.

- As used herein, the term "amplicon" refers to a PCR product amplified from human genomic DNA, containing  
30 the predicted exon.

- As used herein the term "exon" refers to the consensus prediction of the various exon and gene predicting algorithms i.e. a nucleic acid sequence bioinformatically predicted to encode a portion of a  
35 natural protein.

As used herein, the term "peptide" refers to a sequence of amino acids. The sequences referred to as PEPTIDE SEQ ID NOS.: are the predicted peptide sequences that would be translated from one of the exons, or a portion thereof set out in exon SEQ ID NOS.: The codons encoding the peptide are wholly contained within the exon.

As used herein, a "portions" of a defined nucleotide sequence or sequences can be and, preferably, are fragments unique to that sequence or to one or a combination of those sequences. A fragment unique to a nucleic acid molecule is one that is a signature for the larger nucleic acid molecule.

As used herein, the phrase "expression of a probe" and its linguistic variants means that the ORF present within the probe, or its complement, is present within a target mRNA.

As used herein, "stringent conditions" refers to parameters well known to those skilled in the art. When a nucleic acid molecule is said to be hybridisable to another of a given sequence under "stringent conditions" it is meant that it is homologous to the given sequence.

As used herein, the phrase "specific binding pair" intends a pair of molecules that bind to one another with high specificity. Binding pairs are said to exhibit specific binding when they exhibit avidity of at least  $10^7$ , preferably at least  $10^8$ , more preferably at least  $10^9$  liters/mole. Nonlimiting examples of specific binding pairs are: antibody and antigen; biotin and avidin; and biotin and streptavidin.

As used herein with respect to the visual display of annotated genomic sequence, the term "rectangle" means any geometric shape that has at least a first and a second border, wherein the first and second borders each are capable of mapping uniquely to a point of another visual object of the display.

As used herein, a "Mondrian" means a visual display in which a single genomic sequence is annotated with predicted and experimentally confirmed functional information.

5

Brief Description of the Drawings

The present invention is further illustrated with  
10 reference to the following non-limiting figures and examples in which:

FIG. 1 illustrates a process for predicting functional regions from genomic sequence, confirming the functional activity of such regions experimentally, and  
15 associating and displaying the data so obtained in meaningful and useful relationship to the original sequence data;

FIG. 2 further elaborates that portion of the process schematized in FIG. 1 for predicting functional  
20 regions from genomic sequence;

FIG. 3 illustrates a Mondrian visual display;

FIG. 4 presents a Mondrian showing a hypothetical annotated genomic sequence;

FIG. 5 is a histogram showing the distribution of  
25 ORF length and PCR products as obtained, with ORF length shown in black and PCR product length shown in dotted lines;

FIG. 6 is a histogram showing the distribution, among exons predicted according to the methods described,  
30 of expression as measured using simultaneous two color hybridization to a genome-derived single exon microarray. The graph shows the number of sequence-verified products that were either not expressed ("0"), expressed in one or more but not all tested tissues ("1" - "9"), or expressed  
35 in all tissues tested ("10");

FIG. 7 is a pictorial representation of the expression of verified sequences that showed expression with signal intensity greater than 3 in at least one tissue, with: FIG. 7A showing the expression as measured by microarray hybridization in each of the 10 measured tissues, and the expression as measured "bioinformatically" by query of EST, NR and SwissProt databases; with FIG. 7B showing the legend for display of physical expression (ratio) in FIG. 7A; and with FIG. 7C showing the legend for scoring EST hits as depicted in FIG. 7A;

FIG. 8 shows a comparison of normalized CY3 signal intensity for arrayed sequences that were identical to sequences in existing EST, NR and SwissProt databases or that were dissimilar (unknown), where black denotes the signal intensity for all sequence-verified products with a BLAST Expect ("E") value of greater than 1e-30 ( $1 \times 10^{-30}$ ) ("unknown") and a dotted line denotes sequence-verified spots with a BLAST expect ("E") value of less than 1e-30 ( $1 \times 10^{-30}$ ) ("known");

FIG. 9 presents a Mondrian of BAC AC008172 (bases 25,000 to 130,000), containing the carbamyl phosphate synthetase gene (AF154830.1); and

FIG. 10 is a Mondrian of BAC A049839.

25

Methods and Apparatus for Predicting, Confirming,  
Annotating, and Displaying Functional Regions From Genomic  
Sequence Data

FIG. 1 is a flow chart illustrating in broad outline a process for predicting functional regions from genomic sequence, confirming and characterizing the functional activity of such regions experimentally, and then associating and displaying the information so obtained in meaningful and useful relationship to the original

sequence data.

- The initial input into process 10 of the present invention is drawn from one or more databases 100 containing genomic sequence data. Because genomic sequence 5 is usually obtained from subgenomic fragments, the sequence data typically will be stored in a series of records corresponding to these subgenomic sequenced fragments. Some fragments will have been catenated to form larger 10 contiguous sequences ("contigs"); others will not. A finite percentage of sequence data in the database will typically be erroneous, consisting *inter alia* of vector sequence, sequence created from aberrant cloning events, sequence of artificial polylinkers, and sequence that was erroneously read.
- 15 Each sequence record in database 100 will minimally contain as annotation a unique sequence identifier (accession number), and will typically be annotated further to identify the date of accession, species of origin, and depositor. Because database 100 can 20 contain nongenomic sequence, each sequence will typically be annotated further to permit query for genomic sequence. Chromosomal origin, optionally with map location, can also be present. Data can be, and over time increasingly will be, further annotated with additional information, in part 25 through use of the present invention, as described below. Annotation can be present within the data records, in information external to database 100 and linked to the records thereto, or through a combination of the two.
- Databases useful as genomic sequence database 100 30 in the present invention include GenBank, and particularly include several divisions thereof, including the htgs(draft), NT (nucleotide, command line), and NR (nonredundant) divisions. GenBank is produced by the National Institutes of Health and is maintained by the 35 National Center for Biotechnology Information (NCBI).

Databases of genomic sequence from species other than human, such as mouse, rat, *Arabidopsis*, *C. elegans*, *C. briggsii*, *Drosophila*, zebra fish, and other higher eukaryotic organisms will also prove useful as genomic  
5 sequence database 100.

Genomic sequence obtained by query of genomic sequence database 100 is then input into one or more processes 200 for identification of regions therein that are predicted to have a biological function as specified by  
10 the user. Such functions include, but are not limited to, encoding protein, regulating transcription, regulating message transport after transcription into mRNA, regulating message splicing after transcription into mRNA, of regulating message degradation after transcription into  
15 mRNA, and the like. Other functions include directing somatic recombination events, contributing to chromosomal stability or movement, contributing to allelic exclusion or X chromosome inactivation, and the like.

The particular genomic sequence to be input into process 200 will depend upon the function for which relevant sequence is to be identified as well as upon the approach chosen for such identification. Process step 200 can be iterated to identify different functions within a given genomic region. In such case, the input often will  
25 be different for the several iterations.

Sequences predicted to have the requisite function by process 200 are then input into process 300, where a subset of the input sequences suitable for experimental confirmation is identified. Experimental  
30 confirmation can involve physical and/or bioinformatic assay. Where the subsequent experimental assay is bioinformatic, rather than physical, there are fewer constraints on the sequences that can be tested, and in this latter case therefore process 300 can output the  
35 entirety of the input sequence.

The subset of sequences output from process 300 is then used in process 400 for experimental verification and characterization of the function predicted in process 200, which experimental verification can, and often 5 will, include both physical and bioinformatic assay.

Process 500 annotates the sequence data with the functional information obtained in the physical and/or bioinformatic assays of process 400. Such annotation can be done using any technique that usefully relates the 10 functional information to the sequence, as, for example, by incorporating the functional data into the sequence data record itself, by linking records in a hierarchical or relational database, by linking to external databases, by a combination thereof, or by other means well known within 15 the database arts. The data can even be submitted for incorporation into databases maintained by others, such as GenBank, which is maintained by NCBI.

As further noted in FIG. 1, additional annotation can be input into process 500 from external sources 600.

20 The annotated data is then displayed in process 800, either before, concomitantly with, or after optional storage 700 on nontransient media, such as magnetic disk, optical disc, magneto-optical disk, flash memory, or the like.

25 FIG. 1 shows that the experimental data output from process 400 can be used in each preceding step of process 10: e.g., facilitating identification of functional sequences in process 200, facilitating identification of an experimentally suitable subset thereof in process 300, and 30 facilitating creation of physical and/or informational substrates for, and performance of subsequent assay, of functional sequences in process 400.

Information from each step can be passed directly to the succeeding process, or stored in permanent or 35 interim form prior to passage to the succeeding process.

Often, data will be stored after each, or at least a plurality, of such process steps. Any or all process steps can be automated.

FIG. 2 further elaborates the prediction of 5 functional sequence within genomic sequence according to process 200.

Genomic sequence database 100 is first queried 20 for genomic sequence.

The sequence required to be returned by query 20 10 will depend, in the first instance, upon the function to be identified.

For example, genomic sequences that function to encode protein can be identified *inter alia* using gene prediction approaches, comparative sequence analysis 15 approaches, or combinations of the two. In gene prediction analysis, sequence from one genome is input into process 200 where at least one, preferably a plurality, of algorithmic methods are applied to identify putative coding regions. In comparative sequence analysis, by contrast, 20 corresponding, e.g., syntenic, sequence from a plurality of sources, typically a plurality of species, is input into process 200, where at least one, possibly a plurality, of algorithmic methods are applied to compare the sequences and identify regions of least variability.

The exact content of query 20 will also depend 25 upon the database queried. For example, if the database contains both genomic and nongenomic sequence, perhaps derived from multiple species, and the function to be determined is protein coding regions in human genomic 30 sequence, the query will accordingly require that the sequence returned be genomic and derived from humans.

Query 20 can also incorporate criteria that compel return of sequence that meets operative requirements of the subsequent analytical method. Alternatively, or in 35 addition, such operative criteria can be enforced in

subsequent preprocess step 24.

For example, if the function sought to be identified is protein coding, query 20 can incorporate criteria that return from genomic sequence database 100 only those sequences present within contigs sufficiently long as to have obviated substantial fragmentation of any given exon among a plurality of separate sequence fragments.

Such criteria can, for example, consist of a required minimal individual genomic sequence fragment length, such as 10 kb, more typically 20 kb, 30 kb, 40 kb, and preferably 50 kb or more, as well as an optional further or alternative requirement that sequence from any given clone, such as a bacterial artificial chromosome ("BAC"), be presented in no more than a finite maximal number of fragments, such as no more than 20 separate pieces, more typically no more than 15 fragments, even more typically no more than about 10 - 12 fragments.

Results using the present invention have shown that genomic sequence from bacterial artificial chromosomes (BACs) is sufficient for gene prediction analysis according to the present invention if the sequence is at least 50 kb in length, and if additionally the sequence from any given BAC is presented in fewer than 15, and preferably fewer than 10, fragments. Accordingly, query 20 can incorporate a requirement that data accessioned from BAC sequencing be in fewer than 15, preferably fewer than 10, fragments.

An additional criterion that can be incorporated into the query can be the date, or range of dates, of sequence accession. Although the process has been described above as if genomic sequence database 100 were static, it is of course understood that the genomic sequence databases need not be static, and indeed are typically updated on a frequent, even hourly, basis. Thus, as further described in Examples 1 and 2, *infra*, it is

possible to query the database for newly added sequence, either newly added after an absolute date, or newly added relative to a prior analysis performed using the methods and apparatus of the present invention. In this way, the 5 process herein described can incorporate a dynamic, temporal component.

One utility of such temporal limitation is to identify, from newly accessioned genomic sequence, the presence of novel genes, particularly those not previously 10 identified by EST sequencing (or other sequencing efforts that are similarly based upon gene expression). As further described in Example 1, such an approach has shown that newly accessioned human genomic sequence, when analyzed for sequences that function to encode protein, readily 15 identifies genes that are novel over those in existing EST and other expression databases. This makes the methods of the present invention extremely powerful gene discovery tools. And as would be appreciated, such gene discovery can be performed using genomic sequence from species other 20 than human.

If query 20 incorporates multiple criteria, such as above-described, the multiple criteria can be performed as a series of separate queries or as a single query, depending in part upon the query language, the complexity 25 of the query, and other considerations well known in the database arts.

If query 20 returns no genomic sequence meeting the query criteria, the negative result can be reported by process 22, and process 200 (and indeed, entire process 10) 30 ended 23, as shown. Alternatively, or in addition to report and termination of the initial inquiry, a new query 20 can be generated that takes into account the initial negative result.

When query 20 returns sequence meeting the query 35 criteria, the returned sequence is then passed to optional

preprocessing 24, suitable and specific for the desired analytical approach and the particular analytical methods thereof to be used in process 25.

5 Preprocessing 24 can include processes suitable for many approaches and methods thereof, as well as processes specifically suited for the intended subsequent analysis.

10 Preprocessing 24 suitable for most approaches and methods will include elimination of sequence irrelevant to, or that would interfere with, the subsequent analysis. Such sequence includes repetitive sequence, such as Alu repeats and LINE elements, vector sequence, artificial sequence, such as artificial polylinkers, and the like. Such removal can readily be performed by identification and 15 subsequent masking of the undesired sequence.

Identification can be effected by comparing the genomic sequence returned by query 20 with public or private databases containing known repetitive sequence, vector sequence, artificial sequence, and other artifactual sequence. Such comparison can readily be done using programs well known in the art, such as CROSS\_MATCH, or by proprietary sequence comparison programs the engineering of which is well within the skill in the art.

25 Alternatively, or in addition, undesirable, including artifactual, sequence can be identified algorithmically without comparison to external databases and thereafter removed. For example, synthetic polylinker sequence can be identified by an algorithm that identifies a significantly higher than average density of known 30 restriction sites. As another example, vector sequence can be identified by algorithms that identify nucleotide or codon usage at variance with that of the bulk of the genomic sequence.

Once identified, undesired sequence can be 35 removed. Removal can usefully be done by masking the

undesired sequence as, for example, by converting the specific nucleotide references to one that is unrecognized by the subsequent bioinformatic algorithms, such as "X". Alternatively, but at present less preferred, the undesired 5 sequence can be excised from the returned genomic sequence, leaving gaps.

Preprocessing 24 can further include selection from among duplicative sequences of that one sequence of highest quality. Higher quality can be measured as a lower 10 percentage of, fewest number of, or least densely clustered occurrence of ambiguous nucleotides, defined as those nucleotides that are identified in the genomic sequence using symbols indicating ambiguity. Higher quality can also or alternatively be valued by presence in the longest 15 contig.

Preprocessing 24 can, and often will, also include formatting of the data as specifically appropriate for passage to the analytical algorithms of process 25. Such formatting can and typically will include, *inter alia*, 20 addition of a unique sequence identifier, either derived from the original accession number in genomic sequence database 100, or newly applied, and can further include additional annotation. Formatting can include conversion from one to another sequence listing standard, such as 25 conversion to or from FASTA or the like, depending upon the input expected by the subsequent process.

Preprocessing, which can be optional depending upon the function desired to be identified and the informational requirements of the methods for effecting 30 such identification, is followed by sequence processing 25, where sequences with the desired function are identified within the genomic sequence.

As mentioned above, such functions can include, but are not limited to, encoding protein, regulating 35 transcription, regulating message transport after

transcription into mRNA, regulating message splicing after transcription, of regulating message degradation, and the like. Other functions include directing somatic recombination events, contributing to chromosomal stability 5 or movement, contributing to allelic exclusion or X chromosome inactivation, or the like.

The methods of the present invention are particularly useful for gene discovery, that is, for identifying, from genomic sequence, regions that function 10 to encode genes, and in a particularly useful embodiment, for identifying regions that function to encode genes not hitherto identified by expression-based or directed cloning and sequencing. In conjunction with verification using the novel single exon microarrays of the present invention, as 15 further described below, the methods herein described become powerful gene discovery tools.

Accordingly, in a preferred embodiment of the present invention, process 25 is used to identify putative coding regions. Two preferred approaches in process 25 for 20 identifying sequence that encodes putative genes are gene prediction and comparative sequence analysis.

Gene prediction can be performed using any of a number of algorithmic methods, embodied in one or more software programs, that identify open reading frames (ORFs) 25 using a variety of heuristics, such as GRAIL, DICTION, and GENEFINDER. Comparative sequence analysis similarly can be performed using any of a variety of known programs that identify regions with lower sequence variability.

As further described in Example 1, below, gene 30 finding software programs yield a range of results. For the newly accessioned human genomic sequence input in Example 1, for example, GRAIL identified the greatest percentage of genomic sequence as putative coding region, 2% of the data analyzed; GENEFINDER was second, calling 1%; 35 and DICTION yielded the least putative coding region, with

0.8% of genomic sequence called as coding region.

Increased reliability can be obtained when consensus is required among several such methods. Although discussed herein particularly with respect to exon calling,  
5 consensus among methods will in general increase reliability of predicting other functions as well.

Thus, as indicated by query 26, sequence processing 25, optionally with preprocessing 24, can be repeated with a different method, with consensus among such  
10 iterations determined and reported in process 27.

Process 27 compares the several outputs for a given input genomic sequence and identifies consensus among the separately reported results. The consensus itself, as well as the sequence meeting that consensus, is then stored  
15 in process 29a, displayed in process 29b, and/or output to process 300 for subsequent identification of a subset thereof suitable for assay.

Multiple levels of consensus can be calculated and reported by process 27. For example, as further  
20 described in Example 1, *infra*, process 27 can report consensus as between all specific pairs of methods of gene prediction, as consensus among any one or more of the pairs of methods of gene prediction, or as among all of the gene prediction algorithms used. Thus, in Example 1, process 27  
25 reported that GRAIL and GENEFINDER programs agreed on 0.7% of genomic sequence, that GRAIL and DICTION agreed on 0.5% of genomic sequence, and that the three programs together agreed on 0.25% of the data analyzed. Put another way, 0.25% of the genomic sequence was identified by all three  
30 of the programs as containing putative coding region.

Furthermore, consensus can be required among different approaches to identifying a chosen function.

For example, if the function desired to be identified is coding of protein sequence, and a first used  
35 approach to exon calling is gene prediction, the process

can be repeated on the same input sequence, or subset thereof, with another approach, such as comparative sequence analysis. In such a case, where comparative sequence analysis follows gene prediction, the comparison 5 can be performed not only on genomic nucleic acid sequence, but additionally or alternatively can be performed on the predicted amino acid sequence translated from the ORFs prior identified by the gene prediction approach.

Although shown as an iterative process, the 10 multiple analyses required to achieve consensus can be done in series, in parallel, or some combination thereof.

Predicted functional sequence, optionally representing a consensus among a plurality of methods and approaches for determination thereof, is passed to process 15 300 for identification of a subset thereof for functional assay.

In the preferred embodiment of the methods of the present invention, wherein the function sought to be identified is protein coding, process 300 is used to 20 identify a subset thereof suitable for experimental verification by physical and/or bioinformatic approaches.

For example, putative ORFs identified in process 200 can be classified, or binned, bioinformatically into putative genes. This binning can be based *inter alia* upon 25 consideration of the average number of exons/gene in the species chosen for analysis, upon density of exons that have been called on the genomic sequence, and other empirical rules. Thereafter, one or more among the gene-specific ORFs can be chosen for subsequent use in gene 30 expression assay.

Where such subsequent gene expression assay uses amplified nucleic acid, considerations such as desired amplicon length, primer synthesis requirements, putative exon length, sequence GC content, existence of possible 35 secondary structure, and the like can be used to identify

and select those ORFs that appear most likely successfully to amplify. Where subsequent gene expression assay relies upon nucleic acid hybridization, whether or not using amplified product, further considerations involving  
5 hybridization stringency can be applied to identify that subset of sequences that will most readily permit sequence-specific discrimination at a chosen hybridization and wash stringency. One particular such consideration is avoidance of putative exons that span repetitive sequence; such  
10 sequence can hybridize spuriously to nonspecific message, reducing specific signal in the hybridization.

For bioinformatic assay, there are fewer constraints on the sequences that can be tested experimentally, and in this latter case therefore process  
15 300 can output the entirety of the input sequence.

The subset of sequences identified by process 300 as suitable for use in assay is then used in process 400 to create the physical and/or informational substrate for experimental verification of the predictions made in  
20 process 200, and thereafter to assay those substrates.

As mentioned, the methods of the present invention are particularly useful for identifying potential coding regions within genomic sequence. In a preferred embodiment of process 400, therefore, the expression of the  
25 sequences predicted to encode protein is verified. The combination of the predictive and experimental methods provides a powerful gene discovery engine.

Thus, in another aspect, the present invention provides methods and apparatus for verifying the expression  
30 of putative genes identified within genomic sequence. In particular, the invention provides a novel method of verifying gene expression in which expression of predicted ORFs is measured and confirmed using a novel type of nucleic acid microarray, the genome-derived single exon  
35 nucleic acid microarrays of the present invention.

Putative ORFs as predicted by a consensus of gene calling, particularly gene prediction, algorithms in process 200, and as further identified as suitable by process 300, are amplified from genomic DNA using the 5 polymerase chain reaction (PCR). Although PCR is conveniently used, other amplification approaches can also be used.

Amplification schemes can be designed to capture the entirety of each predicted ORF in an amplicon with 10 minimal additional (that is, intronic or intergenic) sequence. Because ORFs predicted from human genomic sequence using the methods of the present invention differ in length, such an approach results in amplicons of varying length.

15 However, most predicted ORFs are shorter than 500 bp in length, and although amplicons of at least about 100 or 200 base pairs can be immobilized as probes on nucleic acid microarrays, early experimental results using the methods of the present invention have suggested that longer 20 amplicons, at least about 400 or 500 base pairs, are more effective. Furthermore, certain advantages derive from application to the microarray of amplicons of defined size.

Therefore, amplification schemes can alternatively, and preferably, be designed to amplify 25 regions of defined size, preferably at least about 300, 400 or 500 bp, centered about each predicted ORF. Such an approach results in a population of amplicons of limited size diversity, but that typically contain intronic and/or intergenic nucleic acid in addition to putative ORF.

30 Conversely, somewhat fewer than 10% of ORFs predicted from human genomic sequence according to the methods of the present invention exceed 500 bp in length. Portions of such extended ORFs, preferably at least about 300, 400 or 500 bp in length, can be amplified. However, it 35 has been discovered that the percentage success at

amplifying pieces of such ORFs is low, and that such putative exons are more effectively amplified when larger fragments, at least about 1000 or 1500 bp, and even as large as 2000 bp are amplified.

- 5       The putative ORFs selected in process 300 are thus input into one or more primer design programs, such as PRIMER3 (available online for use at <http://www-genome.wi.mit.edu/cgi-bin/primer/> ), with a goal of amplifying at least about 500 base pairs of genomic  
10 sequence centered within or about ORFs predicted to be no more than about 500 bp, or at least about 1000 - 1500 bp of genomic sequence for ORFs predicted to exceed 500 bp in length, and the primers synthesized by standard techniques.  
15 Primers with the requisite sequences can be purchased commercially or synthesized by standard techniques.

- Conveniently, a first predetermined sequence can be added commonly to the ORF-specific 5' primer and a second, typically different, predetermined sequence commonly added to each 3' ORF-unique primer. This serves  
20 to immortalize the amplicon, that is, serves to permit further amplification of any amplicon using a single set of primers complementary respectively to the common 5' and common 3' sequence elements. The presence of these "universal" priming sequences further facilitates later  
25 sequence verification, providing a sequence common to all amplicons at which to prime sequencing reactions. The common 5' and 3' sequences further serve to add a cloning site should any of the ORFs warrant further study.

- Such predetermined sequence is usefully at least  
30 about 10, 12 or 15 nt in length, and usually does not exceed about 25 nt in length. The "universal" priming sequences used in the examples presented *infra* were each 16 nt long.

- The genomic DNA to be used as substrate for  
35 amplification will come from the eukaryotic species from

- which the genomic sequence data had originally been obtained, or a closely related species, and can conveniently be prepared by well known techniques from somatic or germline tissue or cultured cells of the
- 5 organism. See, e.g., Short Protocols in Molecular Biology : A Compendium of Methods from Current Protocols in Molecular Biology, Ausubel et al. (eds.), 4<sup>th</sup> edition (April 1999), John Wiley & Sons (ISBN: 047132938X) and Maniatis et al., Molecular Cloning : A Laboratory Manual,
- 10 2<sup>nd</sup> edition (December 1989), Cold Spring Harbor Laboratory Press (ISBN: 0879693096). Many such prepared genomic DNAs are available commercially, with the human genomic DNAs additionally having certification of donor informed consent.
- 15 Although the intronic and intergenic material flanking putative coding regions in the amplicons could potentially interfere with hybridizations during microarray experiments, we have found, surprisingly, that differential expression ratios are not significantly affected. Rather,
- 20 the predominant effect of exon size is to alter the absolute signal intensity, rather than its ratio. Equally surprising, the art had suggested that single exon probes would not provide sufficient signal intensity for high stringency hybridization analyses; we find that such probes
- 25 not only provide adequate signal, but have substantial advantages, as herein described.

After partial purification, as by size exclusion spin column, with or without confirmation as to amplicon quality as by gel electrophoresis, each amplicon (single exon probe) is disposed in an array upon a support substrate.

Methods for creating microarrays by deposition and fixation of nucleic acids onto support substrates are well known in the art (Reviewed by Schena et al., see

35 above).

Typically, the support substrate will be glass, although other materials, such as amorphous or crystalline silicon or plastics. Such plastics include polymethylacrylic, polyethylene, polypropylene, 5 polyacrylate, polymethylmethacrylate, polyvinylchloride, polytetrafluoroethylene, polystyrene, polycarbonate, polyacetal, polysulfone, celluloseacetate, cellulosenitrate, nitrocellulose, or mixtures thereof, can also be used. Typically, the support will be rectangular, 10 although other shapes, particularly circular disks and even spheres, present certain advantages. Particularly advantageous alternatives to glass slides as support substrates for array of nucleic acids are optical discs, as described in WO 98/12559.

15 The amplified nucleic acids can be attached covalently to a surface of the support substrate or, more typically, applied to a derivatized surface in a chaotropic agent that facilitates denaturation and adherence by presumed noncovalent interactions, or some combination 20 thereof.

Robotic spotting devices useful for arraying nucleic acids on support substrates can be constructed using public domain specifications (The MGuide, version 2.0, <http://cmgm.stanford.edu/pbrown/mguide/index.html>), or 25 can conveniently be purchased from commercial sources (MicroArray GenII Spotter and MicroArray GenIII Spotter, Molecular Dynamics, Inc., Sunnyvale, CA). Spotting can also be effected by printing methods, including those using ink jet technology.

30 As is well known in the art, microarrays typically also contain immobilized control nucleic acids. For controls useful in providing measurements of background signal for the genome-derived single exon microarrays of the present invention, a plurality of *E. coli* genes can 35 readily be used. As further described in Example 1, 16 or

32 *E. coli* genes suffice to provide a robust measure of background noise in such microarrays.

- As is well known in the art, the amplified product disposed in arrays on a support substrate to create 5 a nucleic acid microarray can consist entirely of natural nucleotides linked by phosphodiester bonds, or alternatively can include either nonnative nucleotides, alternative internucleotide linkages, or both, so long as complementary binding can be obtained in the hybridization.
- 10 If enzymatic amplification is used to produce the immobilized probes, the amplifying enzyme will impose certain further constraints upon the types of nucleic acid analogs that can be generated.

Although particularly described herein as using 15 high density microarrays constructed on planar substrates, the methods of the present invention for confirming the expression of ORFs predicted from genomic sequence can use any of the known types of microarrays, as herein defined, including lower density planar arrays, and microarrays on 20 nonplanar, nonunitary, distributed substrates.

- For example, gene expression can be confirmed using hybridization to lower density arrays, such as those constructed on membranes, such as nitrocellulose, nylon, and positively-charged derivatized nylon membranes.
- 25 Further, gene expression can also be confirmed using nonplanar, bead-based microarrays such as are described in Brenner et al., *Proc. Natl. Acad. Sci. USA* 97(4):166501670 (2000); U.S. Patent No. 6,057,107; and U.S. Patent No. 5,736,330. In theory, a packed collection of such beads 30 provides in aggregate a higher density of nucleic acid probe than can be achieved with spotting or lithography techniques on a single planar substrate.

Planar microarrays on solid substrates, however, provide certain useful advantages, including high 35 throughput and compatibility with existing readers. For

example, each standard microscope slide can include at least 1000, typically at least 2000, preferably 5000 and upto 10,000 - 50,000 or more nucleic acid probes of discrete sequence. The number of sequences deposited will 5 depend on their required application.

Each putative gene can be represented in the array by a single predicted ORF. Alternatively, genes can be represented by more than one predicted ORF. For purposes of measuring differential splicing, more than one 10 predicted ORF will be provided for a putative gene. And as is well known in the art, each probe of defined sequence, representing a single predicted ORF, can be deposited in a plurality of locations on a single microarray to provide redundancy of signal.

15 The genome-derived single exon microarrays described above differ in several fundamental and advantageous ways from microarrays presently used in the gene expression art, including (1) those created by deposition of mRNA-derived nucleic acids, (2) those created 20 by *in situ* synthesis of oligonucleotide probes, and (3) those constructed from yeast genomic DNA.

Most nucleic acid microarrays that are in use for study of eukaryotic gene expression have as immobilized probes nucleic acids that are derived - either directly or 25 indirectly - from expressed message. As discussed above, it is common, for example, for such microarrays to be derived from cDNA/EST libraries, either from those previously described in the literature, see Lennon et al., or from the *de novo* construction of "problem specific" 30 libraries targeted at a particular biological question, R.S. Thomas et al., *Cancer Res.* (in press). Such microarrays are herein collectively denominated "EST, microarrays".

Such EST microarrays by definition can measure 35 expression only of those genes found in EST libraries,

shown herein to represent only a fraction of expressed genes. Furthermore, such libraries – and thus microarrays based thereupon – are biased by the tissue or cell type of message origin, by the expression levels of the respective 5 genes within the tissues, and by the ability of the message successfully to have been reverse-transcribed and cloned.

Thus, as further discussed in Example 1, the methods of the present invention enable sequences that do not appear in EST or other expression databases to be 10 determined – subsequently arrayed for expression measurements could not, therefore, have been represented as probes on an EST microarray. And as further demonstrated in the examples, *infra*, the remaining population of genes identified from genomic sequence by the methods of the 15 present invention – that is, the one third of sequences that had previously been accessioned in EST or other expression databases – are biased toward genes with higher expression levels.

Representation of a message in an EST and/or cDNA 20 library depends upon the successful reverse transcription, optionally but typically with subsequent successful cloning, of the message. This introduces substantial bias into the population of probes available for arraying in EST microarrays..

In contrast, neither reverse transcription nor 25 cloning is required to produce the probes arrayed on the genome-derived single exon microarrays of the present invention. And although the ultimate deposition of a probe on the genome-derived single exon microarray of the present 30 invention depends upon a successful amplification from genomic material, *a priori* knowledge of the sequence of the desired amplicon affords greater opportunity to recover any given probe sequence recalcitrant to amplification than is afforded by the requirement for successful reverse 35 transcription and cloning of unknown message in EST

approaches.

Thus, the genome-derived single exon microarrays of the present invention present a far greater diversity of probes for measuring gene expression, with far less bias,  
5 than do EST microarrays presently used in the art.

As a further consequence of their ultimate origin from expressed message, the probes in EST microarrays often contain poly-A (or complementary poly-T) stretches derived from the poly-A tail of mature mRNA. These homopolymeric  
10 stretches contribute to cross-hybridization, that is, to a spurious signal occasioned by hybridization to the homopolymeric tail of a labeled cDNA that lacks sequence homology to the gene-specific portion of the probe.

In contrast, the probes arrayed in the genome-derived single exon microarrays of the present invention lack homopolymeric stretches derived from message polyadenylation, and thus can provide more specific signal.  
15 Typically, at least about 50, 60 or 75% of the probes on the genome-derived single exon microarrays of the present invention lack homopolymeric regions consisting of A or T, where a homopolymeric region is defined for purposes herein as stretches of 25 or more, typically 30 or more, identical nucleotides.  
20

A further distinction, which also affects the specificity of hybridization, is occasioned by the typical derivation of EST microarray probes from cloned material.  
25 Because much of the probe material disposed as probes on EST microarrays is excised or amplified from plasmid, phage, or phagemid vectors, EST microarrays typically include a fair amount of vector sequence, more so when the probes are amplified, rather than excised, from the vector.  
30

In contrast, the vast majority of probes in the genome-derived single exon microarrays of the present invention contain no prokaryotic or bacteriophage vector  
35 sequence, having been amplified directly or indirectly from

genomic DNA. Typically, therefore, at least about 50, 60, 70 or 80% or more of individual exon-including probes disposed on a genome-derived single exon microarray of the present invention lack vector sequence, and particularly 5 lack sequences drawn from plasmids and bacteriophage. Preferably, at least about 85, 90 or more than 90% of exon-including probes in the genome-derived single exon microarray of the present invention lack vector sequence. With attention to removal of vector sequences through 10 preprocessing 24, percentages of vector-free exon-including probes can be as high as 95 - 99%. The substantial absence of vector sequence from the genome-derived single exon microarrays of the present invention results in greater specificity during hybridization, since spurious cross- 15 hybridization to a probe vector sequence is reduced.

As a further consequence of excision or amplification of probes from vectors in construction of EST microarrays, the probes arrayed thereon often contain artificial sequence, derived from vector polylinker 20 multiple cloning sites, at both 5' and 3' ends. The probes disposed upon the genome-derived single exon microarrays need have no such artificial sequence appended thereto.

As mentioned above, however, the ORF-specific primers used to amplify putative ORFs can include 25 artificial sequences, typically 5' to the ORF-specific primer sequence, useful for "universal" (that is, independent of ORF sequence) priming of subsequent amplification or sequencing reactions. When such "universal" 5' and/or 3' priming sequences are appended to 30 the amplification primers, the probes disposed upon the genome-derived single exon microarray will include artificial sequence similar to that found in EST microarrays. However, the genome-derived single exon microarray of the present invention can be made without 35 such sequences, and if so constructed, presents an even

smaller amount of nonspecific sequence that would contribute to nonspecific hybridization.

Yet another consequence of typical use of cloned material as probes in EST microarrays is that such 5 microarrays contain probes that result from cloning artifacts, such as chimeric molecules containing coding region of two separate genes. Derived from genomic material, typically not thereafter cloned, the probes of the genome-derived single exon microarrays of the present 10 invention lack such cloning artifacts, and thus provide greater specificity of signal in gene expression measurements.

A further consequence of the cloned origin of probes on many EST microarrays is that the individual 15 probes often have disparate sizes, which can cause the optimal hybridization stringency to vary among probes on a single microarray. In contrast, as discussed above, the probes arrayed on the genome-derived single exon microarrays of the present invention can readily be 20 designed to have a narrow distribution in sizes, with the range of probe sizes no greater than about 10% of the average size, typically no greater than about 5% of the average probe size.

Because of their origin from fully- or partially-spliced message, probes disposed upon EST arrays will often 25 include multiple exons. The percentage of such exon-spanning probes in an EST microarray can be calculated, on average, based upon the predicted number of exons/gene for the given species and the average length of the immobilized 30 probes. For human genes, the near-complete sequence of human chromosome 22, Dunham *et al.*, *Nature* 402(6761):489-95 (1999), predicts that human genes average 5.5 exons/gene. Even with probes of 200 - 500 bp, the vast majority of 35 human EST microarray probes include more than one exon.

In contrast, by virtue of their origin from

algorithmically identified ORFs in genomic sequence, the probes in the genome-derived single exon microarrays of the present invention can consist of individual exons. Thus, in contrast to EST microarrays, at least about 50, 60, 70, 5 75, 80, 85, 95 or 99% of probes deposited in the genome-derived microarray of the present invention consist of, or include, no more than one predicted ORF.

This provides the ability, not readily achieved using EST microarrays, to use the genome-derived single 10 exon microarrays of the present invention to measure tissue-specific expression of individual exons, which in turn allows differential splicing events to be detected and characterized, and in particular, allows the correlation of differential splicing to tissue-specific expression 15 patterns.

Furthermore, the exons that are represented in EST microarrays are often biased toward the 3' or 5' end of their respective genes, since sequencing strategies used for EST identification are so biased. In contrast, no such 20 3' or 5' bias necessarily inheres in the selection of exons for disposition on the genome-derived single exon microarrays of the present invention.

Conversely, the probes provided on the genome-derived single exon microarrays of the present invention 25 typically, but need not necessarily, include intronic and/or intergenic sequence that is absent from EST microarrays, which are derived from mature mRNA. Typically, at least about 50, 60, 70, 80 or 90% of the exon-including probes on the genome-derived single exon 30 microarrays of the present invention include sequence drawn from noncoding regions. As discussed above, the additional presence of noncoding region does not significantly interfere with measurement of gene expression, and provides the additional opportunity to assay prespliced RNA, and 35 thus measure such phenomena such as nuclear export control.

The genome-derived single exon microarrays of the present invention are also quite different from *in situ* synthesis microarrays, where probe size is severely constrained by inadequacies in the photolithographic synthesis process.

- Typically, probes arrayed on *in situ* synthesis microarrays are limited to a maximum of about 25 bp. As a well known consequence, hybridization to such chips must be performed at low stringency. In order, therefore, to
- 10 achieve unambiguous sequence-specific hybridization results, the *in situ* synthesis microarray requires substantial redundancy, with concomitant programmed arraying for each probe of probe analogues with altered (i.e., mismatched) sequence.
- 15 In contrast, the longer probe length of the genome-derived single exon microarrays of the present invention allows much higher stringency hybridization and wash. Typically, therefore, exon-including probes on the genome-derived single exon microarrays of the present
- 20 invention average at least about 100, 200, 300, 400 or 500 bp in length. By obviating the need for substantial probe redundancy, this approach permits a higher density of probes for discrete exons or genes to be arrayed on the microarrays of the present invention than can be achieved
- 25 for *in situ* synthesis microarrays.

A further distinction is that the probes in *in situ* synthesis microarrays typically are covalently linked to the substrate surface. In contrast, the probes disposed on the genome-derived microarray of the present invention

30 typically are, but need not necessarily be, bound noncovalently to the substrate.

Furthermore, the short probe size on *in situ* microarrays causes large percentage differences in the melting temperature of probes hybridized to their

35 complementary target sequence, and thus causes large

percentage differences in the theoretically optimum stringency across the array as a whole.

In contrast, the larger probe size in the microarrays of the present invention create lower

5 percentage differences in melting temperature across the range of arrayed probes.

A further significant advantage of the microarrays of the present invention over *in situ* synthesized arrays is that the quality of each individual 10 probe can be confirmed before deposition. In contrast, the quality of probes cannot be assessed on a probe-by-probe basis for the *in situ* synthesized microarrays presently being used.

The genome-derived single exon microarrays of the 15 present invention are also distinguished over, and present substantial benefits over, the genome-derived microarrays from lower eukaryotes such as yeast. Lashkari et al., *Proc. Natl. Acad. Sci. USA* 94:13057-13062 (1997).

Only about 220 - 250 of the 6100 or so nuclear 20 genes in *Saccharomyces cerevisiae* - that is, only about 4 - 5% - have standard, spliceosomal, introns, Lopez et al., *Nucl. Acids Res.* 28:85-86 (2000); Spingola et al., *RNA* 5(2):221-34 (1999). Furthermore, the entire yeast genome has already been sequenced. These two facts permit the 25 ready amplification and disposition of single-ORF amplicons on such microarray without the requirement for antecedent use of gene prediction and/or comparative sequence analyses.

Thus, a significant aspect of the present 30 invention is the ability to identify and to confirm expression of predicted coding regions in genomic sequence drawn from eukaryotic organisms that have a higher percentage of genes having introns than do yeast such as *Saccharomyces cerevisiae*, particularly in genomic sequence 35 drawn from eukaryotes in which at least about 10, 20 or 50%

of protein-encoding genes have introns. In preferred embodiments, the methods and apparatus of the present invention are used to identify and confirm expression of novel genes from genomic sequence of eukaryotes in which  
5 the average number of introns per gene is at least about one, two or three or more.

After the physical substrate is prepared, experimental verification of predicted function is performed.

10 In a preferred embodiment of the present invention, where the function sought to be identified in genomic sequence is protein coding, experimental verification is performed by measuring expression of the putative ORFs, typically through nucleic acid hybridization  
15 experiments, and in particularly preferred embodiments, through hybridization to genome-derived single exon microarrays prepared as above-described.

Expression is conveniently measured and expressed for each probe in the microarray as a ratio of the  
20 expression measured concurrently in a plurality of mRNA sources, according to techniques well known in the microarray art, *Reviewed in Schena et al.*, and as further described in Example 2, below. The mRNA source for the reference against which specific expression is measured can  
25 be drawn from a homogeneous mRNA source, such as a single cultured cell-type, or alternatively can be heterogeneous, as from a pool of mRNA derived from multiple tissues and/or cell types, as further described in Example 2, *infra*.

mRNA can be prepared by standard techniques, see  
30 Ausubel et al. and Maniatis et al., or purchased commercially. The mRNA is then typically reverse-transcribed in the presence of labeled nucleotides: the index source (that in which expression is desired to be measured) is reverse transcribed in the presence of  
35 nucleotides labeled with a first label, typically a

fluorophore (fluorochrome; fluor; fluorescent dye); the reference source is reverse transcribed in the presence of a second label, typically a fluorophore, typically fluorometrically-distinguishable from the first label. As 5 further described in Example 2, *infra*, Cy3 and Cy5 dyes prove particularly useful in these methods. After partial purification of the index and reference targets, hybridization to the probe array is conducted according to standard techniques, typically under a coverslip.

10 After wash, microarrays are conveniently scanned using a commercial microarray scanning device, such as a Gen3 Scanner (Molecular Dynamics, Sunnyvale, CA). Data on expression is then passed, with or without interim storage, to process 500, where the results for each probe are 15 related to the original sequence.

Often, hybridization of target material to the genome-derived single exon microarray will identify certain of the probes thereon as of particular interest. Thus, it is often desirable that the user be able readily to obtain 20 sufficient quantities of an individual probe, either for subsequent arrayed deposition upon an additional support substrate, often as part of a microarray having a plurality of probes so identified, or alternatively or additionally as a solitary solid-phase or solution-phase probe, for 25 further use.

Thus, in another aspect, the present invention provides compositions and kits for the ready production of nucleic acids identical in sequence to, or substantially identical in sequence to, probes on the genome-derived 30 single exon microarrays of the present invention.

In this aspect, a small quantity of each probe is disposed, typically without attachment to substrate, in a spatially-addressable ordered set, typically one per well of a microtiter dish. Although a 96 well microtiter plate 35 can be used, greater efficiency is obtained using higher

density arrays, such as are provided by microtiter plates having 384, 864, 1536, 3456, 6144, or 9600 wells, and although microtiter plates having physical depressions (wells) are conveniently used, any device that permits 5 addressable withdrawal of reagent from fluidly-noncommunicating areas can be used.

In this aspect of the invention, therefore, a fluidly noncommunicating addressable ordered set of individual probes, corresponding to those on a genome-10 derived single exon microarray, is provided, with each probe in sufficient quantity to permit amplification, such as by PCR. As earlier mentioned, the ORF-specific 5' primers used for genomic amplification can have a first common sequence added thereto, and the ORF-specific 3' 15 primers used for genomic amplification can have a second, different, common sequence added thereto, thus permitting, in this preferred embodiment, the use of a single set of 5' and 3' primers to amplify any one of the probes from the amplifiable ordered set.

20 Each discrete amplifiable probe can also be packaged with amplification primers, solutes, buffers, etc., and can be provided in dry (e.g., lyophilized) form or wet, in the latter case typically with addition of agents that retard evaporation.

25 In another aspect of the present invention, a genome-derived single-exon microarray is packaged together with such an ordered set of amplifiable probes corresponding to the probes, or one or more subsets of probes, thereon. In alternative embodiments, the ordered 30 set of amplifiable probes is packaged separately from the genome-derived single exon microarray.

In some embodiments, the microarray and/or ordered probe set are further packaged with recordable media that provide probe identification and addressing 35 information, and that can additionally contain annotation

information, such as gene expression data. Such recordable media can be packaged with the microarray, with the ordered probe set, or with both.

If the microarray is constructed on a substrate 5 that incorporates recordable media, such as is described in international patent application no. WO 98/12559, then separate packaging of the genome-derived single exon microarray and the bioinformatic information is not required.

10 The amount of amplifiable probe material should be sufficient to permit at least one amplification sufficient for subsequent hybridization assay.

Although the use of high density genome-derived microarrays on solid planar substrates is presently a 15 preferred approach for the physical confirmation and characterization of the expression of sequences predicted to encode protein, other types of microarrays (as herein defined) can also be used.

Furthermore, as earlier mentioned, experimental 20 verification of the function predicted from genomic sequence in process 200 can be bioinformatic, rather than, or additional to, physical verification.

For example, where the function desired to be identified is protein coding, the predicted ORFs can be 25 compared bioinformatically to sequences known or suspected of being expressed.

Thus, the sequences output from process 300 (or process 200), can be used to query expression databases, such as EST databases, SNP ("single nucleotide 30 polymorphism") databases, known cDNA and mRNA sequences, SAGE ("serial analysis of gene expression") databases, and more generalized sequence databases that allow query for expressed sequences. Such query can be done by any sequence query algorithm, such as BLAST ("basic local 35 alignment search tool"). The results of such query -

including information on identical sequences and information on nonidentical sequences that have diffuse or focal regions of sequence homology to the query sequence – can then be passed directly to process 500, or used to 5 inform analyses subsequently undertaken in process 200, process 300, or process 400.

Experimental data, whether obtained by physical or bioinformatic assay in process 400, is passed to process 500 where it is usefully related to the sequence data 10 itself, a process colloquially termed "annotation". Such annotation can be done using any technique that usefully relates the functional information to the sequence, as, for example, by incorporating the functional data into the record itself, by linking records in a hierarchical or 15 relational database, by linking to external databases, or by a combination thereof. Such database techniques are well within the skill in the art.

The annotated sequence data can be stored locally, uploaded to genomic sequence database 100, and/or 20 displayed 800.

The methods and apparatus of the present invention rapidly produce functional information from genomic sequence. Coupled with the escalating pace at which sequence now accumulates, the rapid pace of sequence 25 annotation produces a need for methods of displaying the information in meaningful ways.

FIG. 3 shows visual display 80 presenting a single genomic sequence annotated according to the present invention. Because of its nominal resemblance to artistic 30 works of Piet Mondrian, visual display 80 is alternatively described herein as a "Mondrian".

Each of the visual elements of display 80 is aligned with respect to the genomic sequence being annotated (hereinafter, the "annotated sequence"). Given 35 the number of nucleotides typically represented in an

annotated sequence, representation of individual nucleotides would rarely be readable in hard copy output of display 80. Typically, therefore, the annotated sequence is schematized as rectangle 89, extending from the left border of display 80 to its right border. By convention herein, the left border of rectangle 89 represents the first nucleotide of the sequence and the right border of rectangle 89 represents the last nucleotide of the sequence.

As further discussed below, however, the Mondrian visual display of annotated sequence can serve as a convenient graphical user interface for computerized representation, analysis, and query of information stored electronically. For such use, the individual nucleotides can conveniently be linked to the X axis coordinate of rectangle 89. This permits the annotated sequence at any point within rectangle 89 readily to be viewed, either automatically - for example, by time-delayed appearance of a small overlaid window upon movement of a cursor or other pointer over rectangle 89 - or through user intervention, as by clicking a mouse or other pointing device at a point in rectangle 89.

Visual display 80 is generated after user specification of the genomic sequence to be displayed. Such specification can consist of or include an accession number for a single clone (e.g., a single BAC accessioned into GenBank), wherein the starting and stopping nucleotides are thus absolutely identified, or alternatively can consist of or include an anchor or fulcrum point about which a chosen range of sequence is anchored, thus providing relative endpoints for the sequence to be displayed. For example, the user can anchor such a range about a given chromosomal map location, gene name, or even a sequence returned by query for similarity or identity to an input query sequence. When visual

display 80 is used as a graphical user interface to computerized data, additional control over the first and last displayed nucleotide will typically be dynamically selectable, as by use of standard zooming and/or selection tools.

Field 81 of visual display 80 is used to present the output from process 200, that is, to present the bioinformatic prediction of those sequences having the desired function within the genomic sequence. Functional sequences are typically indicated by at least one rectangle 83 (83a, 83b, 83c), the left and right borders of which respectively indicate, by their X-axis coordinates, the starting and ending nucleotides of the region predicted to have function.

Where a single bioinformatic method or approach identifies a plurality of regions having the desired function, a plurality of rectangles 83 is disposed horizontally in field 81. Where multiple methods and/or approaches are used to identify function, each such method and/or approach can be represented by its own series of horizontally disposed rectangles 83, each such horizontally disposed series of rectangles offset vertically from those representing the results of the other methods and approaches.

Thus, rectangles 83a in FIG. 3 represent the functional predictions of a first method of a first approach for predicting function, rectangles 83b represent the functional predictions of a second method and/or second approach for predicting that function, and rectangles 83c represent the predictions of a third method and/or approach.

Where the function desired to be identified is protein coding, field 81 is used to present the bioinformatic prediction of sequences encoding protein. For example, rectangles 83a can represent the results from

GRAIL or GRAIL II, rectangles 83b can represent the results from GENEFINDER, and rectangles 83c can represent the results from DICTION.

Optionally, and preferably, rectangles 83  
5 collectively representing predictions of a single method and/or approach are identically colored and/or textured, and are distinguishable from the color and/or texture used for a different method and/or approach.

Alternatively, or in addition, the color, hue,  
10 density, or texture of rectangles 83 can be used further to report a measure of the bioinformatic reliability of the prediction. For example, many gene prediction programs will report a measure of the reliability of prediction. Thus, increasing degrees of such reliability can be  
15 indicated, e.g., by increasing density of shading. Where display 80 is used as a graphical user interface, such measures of reliability, and indeed all other results output by the program, can additionally or alternatively be made accessible through linkage from individual rectangles  
20 83, as by time-delayed window ("tool tip" window), or by pointer (e.g., mouse)-activated link.

As earlier described, increased predictive reliability can be achieved by requiring consensus among methods and/or approaches to determining function. Thus,  
25 field 81 can include a horizontal series of rectangles 83 that indicate one or more degrees of consensus in predictions of function.

Although FIG. 3 shows three series of horizontally disposed rectangles in field 81, display 80  
30 can include as few as one such series of rectangles and as many as can discriminably be displayed, depending upon the number of methods and/or approaches used to predict a given function.

Furthermore, field 81 can be used to show  
35 predictions of a plurality of different functions.

However, the increased visual complexity occasioned by such display makes more useful the ability of the user to select a single function for display. When display 80 is used as a graphical user interface for computer query and analysis, 5 such function can usefully be indicated and user-selectable, as by a series of graphical buttons or tabs (not shown in FIG. 3).

Rectangle 89 is shown in FIG. 3 as including interposed rectangle 84. Rectangle 84 represents the 10 portion of annotated sequence for which predicted functional information has been assayed physically, with the starting and ending nucleotides of the assayed material indicated by the X axis coordinates of the left and right borders of rectangle 84. Rectangle 85, with optional 15 inclusive circles 86 (86a, 86b, and 86c) displays the results of such physical assay.

Although a single rectangle 84 is shown in FIG. 3, physical assay is not limited to just one region of annotated genomic sequence. It is expected that an 20 increasing percentage of regions predicted to have function by process 200 will be assayed physically, and that display 80 will accordingly, for any given genomic sequence, have an increasing number of rectangles 84 and 85, representing an increased density of sequence annotation.

25 Where the function desired to be identified is protein coding, rectangle 84 identifies the sequence of the probe used to measure expression. In embodiments of the present invention where expression is measured using genome-derived single exon microarrays, rectangle 84 30 identifies the sequence included within the probe immobilized on the support surface of the microarray. As noted *supra*, such probe will often include a small amount of additional, synthetic, material incorporated during amplification and designed to permit reamplification of the 35 probe, which sequence is typically not shown in display 80.

- Rectangle 87 is used to present the results of bioinformatic assay of the genomic sequence. For example, where the function desired to be identified is protein coding, process 400 can include bioinformatic query of expression databases with the sequences predicted in process 200 to encode exons. And as earlier discussed, because bioinformatic assay presents fewer constraints than does physical assay, often the entire output of process 200 can be used for such assay, without further subsetting thereof by process 300. Therefore, rectangle 87 typically need not have separate indicators therein of regions submitted for bioinformatic assay; that is, rectangle 87 typically need not have regions therein analogous to rectangles 84 within rectangle 89.
- Rectangle 87 as shown in FIG. 3 includes smaller rectangles 880 and 88. Rectangles 880 indicate regions that returned a positive result in the bioinformatic assay, with rectangles 88 representing regions that did not return such positive results. Where the function desired to be predicted and displayed is protein coding, rectangles 880 indicate regions of the predicted exons that identify sequence with significant similarity in expression databases, such as EST, SNP, SAGE databases, with rectangles 88 indicating genes novel over those identified in existing expression data bases.

Rectangles 880 can further indicate, through color, shading, texture, or the like, additional information obtained from bioinformatic assay.

For example, where the function assayed and displayed is protein coding, the degree of shading of rectangles 880 can be used to represent the degree of sequence similarity found upon query of expression databases. The number of levels of discrimination can be as few as two (identity, and similarity, where similarity has a user-selectable lower threshold). Alternatively, as

many different levels of discrimination can be indicated as can visually be discriminated.

Where display 80 is used as a graphical user interface, rectangles 880 can additionally provide links directly to the sequences identified by the query of expression databases, and/or statistical summaries thereof. As with each of the precedingly-discussed uses of display 80 as a graphical user interface, it should be understood that the information accessed via display 80 need not be resident on the computer presenting such display, which often will be serving as a client, with the linked information resident on one or more remotely located servers.

Rectangle 85 displays the results of physical assay of the sequence delimited by its left and right borders.

Rectangle 85 can consist of a single rectangle, thus indicating a single assay, or alternatively, and increasingly typically, will consist of a series of rectangles (85a, 85b, 85c) indicating separate physical assays of the same sequence.

Where the function assayed is gene expression, and where gene expression is assayed as herein described using simultaneous two-color fluorescent detection of hybridization to genome-derived single exon microarrays, individual rectangles 85 can be colored to indicate the degree of expression relative to control. Conveniently, shades of green can be used to depict expression in the sample over control values, and shades of red used to depict expression less than control, corresponding to the spectra of the Cy3 and Cy5 dyes conventionally used for respective labeling thereof. Additional functional information can be provided in the form of circles 86 (86a, 86b, 86c), where the diameter of the circle can be used to indicate expression intensity. As discussed *infra*, such

relative expression (expression ratios) and absolute expression (signal intensity) can be expressed using normalized values.

Where display 80 is used as a graphical user interface, rectangle 85 can be used as a link to further information about the assay. For example, where the assay is one for gene expression, each rectangle 85 can be used to link to information about the source of the hybridized mRNA, the identity of the control, raw or processed data from the microarray scan, or the like.

FIG. 4 is rendition of display 80 representing gene prediction and gene expression for a hypothetical BAC, showing conventions used in the Examples presented *infra*. BAC sequence ("Chip seq.") 89 is presented, with the physically assayed region thereof (corresponding to rectangle 84 in FIG. 3) shown in white. Algorithmic gene predictions are shown in field 81, with predictions by GRAIL shown, predictions by GENEFINDER, and predictions by DICTION shown. Within rectangle 87, regions of sequence that, when used to query expression databases, return identical or similar sequences ("EST hit") are shown as white rectangles (corresponding to rectangles 880 in FIG. 3), gray indicates low homology, and black indicates unknowns (where black and gray would correspond to rectangles 88 in FIG. 3).

Although FIGS. 3 and 4 show a single stretch of sequence, uninterrupted from left to right, longer sequences are usefully represented by vertical stacking of such individual Mondrians, as shown in FIGS. 9 and 10.

30

#### Single Exon Probes Useful For Measuring Gene Expression

The methods and apparatus of the present invention rapidly produce functional information from genomic sequence. Where the function to be identified is

protein coding, the methods and apparatus of the present invention rapidly identify and confirm the expression of portions of genomic sequence that function to encode protein. As a direct result, the methods and apparatus of 5 the present invention rapidly yield large numbers of single-exon nucleic acid probes, the majority from previously unknown genes, each of which is useful for measuring and/or surveying expression of a specific gene in one or more tissues or cell types.

10 It is, therefore, another aspect of the present invention to provide genome-derived single exon nucleic acid probes useful for gene expression analysis, and particularly for gene expression analysis by microarray.

15 Using the methods and genome-derived single-exon microarrays of the present invention, we have for example readily identified a large number of unique ORFs from human genomic sequence. Using single exon probes that encompass these ORFs, we have demonstrated, through microarray hybridization analysis, the expression of 12,821 of these 20 ORFs in brain.

As would immediately be appreciated by one of skill in the art, each single exon probe having demonstrable expression in brain is currently available for use in measuring the level of its ORF's expression in 25 brain.

Diseases of the brain and nervous system are a significant cause of human morbidity and mortality. Increasingly, genetic factors are being found that contribute to predisposition, onset, and/or aggressiveness 30 of most, if not all, of these diseases. Although mutations in single genes have been identified as causative for some diseases of the brain and nervous system, for the most part these disorders are believed to have polygenic etiologies.

For example, over the past few decades 35 Alzheimer's disease (AD), once considered a rare disorder,

has become recognized as a major public health problem; over 4,000,000 people in the United States are now estimated to suffer with various stages of this progressive, degenerative brain disorder.

- 5        Although there is no agreement on the exact incidence or prevalence of Alzheimer's disease, in part due to varying diagnostic criteria and difficulties of differential diagnosis among dementias, the studies are consistent in pointing to an exponential rise in prevalence  
10      of this disease with age. After age 65, the percentage of affected people approximately doubles with every decade of life, regardless of definition. Among people age 85 or older, studies suggest that 25 to 35 percent have dementia, including Alzheimer's disease; one study reports that 47.2  
15      percent of people over age 85 have Alzheimer's disease, exclusive of other dementias.

Alzheimer's disease progressively destroys memory, reason, judgment, language, and, eventually, the ability to carry out even the simplest of tasks. Anatomic  
20      changes associated with Alzheimer's disease begin in the entorhinal cortex, proceed to the hippocampus, and then gradually spread to other regions, particularly the cerebral cortex. Chief among such anatomic changes are the presence of characteristic extracellular plaques and  
25      internal neurofibrillary tangles.

Alzheimer's disease has been suspected to have a multifactorial genetic etiological component for almost half a century. Sjogren et al., *Acta Psychiat. Neurol. Scand.* 82(suppl.): 1-152 (1952).

- 30       At least four genes have been identified to date that contribute to development of Alzheimer's disease: AD1 is caused by mutations in the amyloid precursor gene (APP); AD2 is associated with the APOE4 allele on chromosome 19; AD3 is caused by mutation in a chromosome 14 gene encoding  
35      a 7-transmembrane domain protein, presenilin-1 (PSEN1), and

AD4 is caused by mutation in a gene on chromosome 1 that encodes a similar 7-transmembrane domain protein, presenilin-2 (PSEN2).

There is strong evidence, however, for  
5 additional, as yet uncharacterized, AD loci on other chromosomes.

For example, Daw et al., Am. J. Hum. Genet. 66: 196-204 (2000), estimated the number of additional quantitative trait loci (QTLs) and their contribution to  
10 the variance in age at onset of AD, and reported that 4 loci make a contribution to the variance in age at onset of late-onset AD similar to or greater in magnitude than that made by apoE, with one locus making a contribution several times greater than that of  
15 apoE. These results suggest that several genes not yet localized may play a larger role than does apoE in late-onset AD.

In accord, three groups recently announced the possible existence of an AD susceptibility gene on  
20 chromosome 10. Bertram et al., Science 290(5500):2302-2303 (2000); Ertekin-Taner et al., Science 290(5500):2303-2304 (2000); and Myers et al., Science 290(5500):2304-23055 (2000).

As another example, multiple sclerosis (MS)  
25 affects about 350,000 Americans, with approximately 200 new cases diagnosed each week, with an estimated annual monetary cost in the U.S. alone of \$2.5 billion.

Clinically, MS is an unpredictable disorder, with symptoms, presentation and course falling broadly into one  
30 of several clinical patterns. In relapsing-remitting (RR) MS, the disease first manifests as a series of attacks followed by complete or partial remissions, with symptoms returning later after a period of stability. In primary-progressive (PP) MS, there is a gradual clinical decline  
35 with no distinct remissions, although there may be

temporary plateaus or minor relief from symptoms.

Secondary-progressive (SP) MS begins with a relapsing-remitting course followed by a later primary-progressive course. Rarely, patients may have a progressive-relapsing

5 (PR) course in which the disease takes a progressive path punctuated by acute attacks. PP, SP, and PR MS are sometimes lumped together and called chronic progressive MS. The waxing and waning course characteristic of RR, SP and PR MS makes differential diagnosis difficult.

10 Anatomically, MS attacks are associated with focal inflammation in areas of the white matter of the central nervous system (CNS), accompanied or followed by demyelination in these areas, termed plaques. Destruction of the myelin sheath slows or blocks neurological

15 transmission, leading to diminished or lost function. Clinical manifestations depend upon the location of the plaques and severity of demyelination, and range from fatigue, the most common symptom of MS, to visual impairment, due to inflammation of the optic nerve, termed  
20 optic neuritis, to numbness and paresthesias, to focal muscular weakness, ataxia, and bladder incontinence.

Increasing evidence suggests that genotype contributes to susceptibility to MS.

As early as 1965, McAlpine, in Multiple  
25 Sclerosis: A Reappraisal (McAlpine, ed.), Williams and Wilkins Co. pp. 61-74 (1965), concluded that the risk to a first-degree relative of a patient with multiple sclerosis is at least 15 times that for a member of the general population, but could discern no definite genetic pattern  
30 of inheritance.

Subsequently, many studies associated MS with HLA (MHC) haplotype. Haines et al., Hum. Molec. Genet. 7:1229-1234 (1998), studying a data set of 98 multiplex MS families, confirmed earlier reports that genetic linkage to  
35 the MHC can be explained by association with the HLA-DR2

allele, but suggested that MHC association explains only between 17% and 62% of the genetic etiology of MS.

From a review of genomic screens, Dyment et al., Hum. Molec. Genet. 6: 1693-1698 (1997), concluded that a number of genes with interacting effects are likely and that no single region has a major influence on familial risk. Chataway et al., Brain 121: 1869-1887 (1998), reporting a follow-up on U.K. studies using a systematic genome screen to determine the genetic basis of MS, stated that a gene of major effect had been excluded from 95% of the genome and one with a moderate role from 65%, results thus suggesting that multiple sclerosis depends on independent or epistatic effects of several genes, each with small individual effects, rather than a very few genes of major biologic importance.

As a yet further example, schizophrenia has long been recognized to have complex, likely polygenic, genetic contributions.

Schizophrenia is a common psychiatric disorder, occurring in 1 to 1.5 percent of the population worldwide, and is characterized by variable constellations of symptoms drawn from a universe of behavioral abnormalities. Although there are accepted alternative diagnostic criteria, primary criteria for diagnosis require two or more of the following, each present for a significant portion of time during a 1-month period (or less if successfully treated): (1) delusions; (2) hallucinations; (3) disorganized speech (e.g., frequent derailment or incoherence); (4) grossly disorganized or catatonic behavior; (5) negative symptoms, i.e., affective flattening, alogia, or avolition. (Diagnostic and Statistic Manual of Mental Disorders DSM-IV-TR, American Psychiatric Association (2000)). Only one such symptom is required if delusions are bizarre or hallucinations consist of a voice keeping up a running commentary on the

person's behavior or thoughts, or consist of two or more voices conversing with each other.

Three-quarters of persons with schizophrenia develop the disease between 16 and 25 years of age: onset is uncommon after age 30, rare after age 40. In the 16 to 25 year old age group, schizophrenia affects more men than women; in the 25-30 year old group, the incidence is higher in women than in men. Studies have shown that some persons with schizophrenia recover completely, and many others improve to the point where they can live independently, often with the maintenance of drug therapy. However, approximately 15 percent of people with schizophrenia respond only moderately to medication and require extensive support throughout their lives, while another 15 percent simply do not respond to existing treatment.

Schizophrenia has long been known to have a significant genetic component. Studies have consistently demonstrated that the risk to relatives of a proband with schizophrenia is higher than the risk to relatives of controls. Moldin, in Genetics and Mental Disorders: Report of the NIMH Genetics Workgroup (NIH publication 98-4268, (1998), reviewed family and twin studies published between 1920 and 1987 and found the recurrence risk ratios to be 48 for monozygotic twins, 11 for first-degree relatives, 4.25 for second-degree relatives, and 2 for third-degree relatives. He also found that concordance rates for monozygotic twins averaged 46%, even when reared in different families, whereas the concordance rates for dizygotic twins averaged only 14%. The prevalence of schizophrenia is known to be higher in biologic than in adoptive relatives of schizophrenic adoptees.

The mode of inheritance is unclear, however. Susceptibility has been mapped to many loci, including chromosomes 1q21-q22, 5, 6p23, 8p22-p21, 11q, 13q14-q21, 35 13q32, 15q15, 15q14, 18p, and 22q11. Chromosome

19 has also been implicated in schizophrenia, at 2 different sites, as have sites on the X chromosome. Wei et al., Nature Genet. 25:376-377 (2000) report more specifically that the NOTCH4 locus is associated with 5 susceptibility to schizophrenia.

In general, however, it is believed that development of schizophrenia involves multiple loci.

For example, Williams et al., Hum. Molec. Genet. 8:1729-1739 (1999) undertook a systematic search for 10 linkage in 196 affected sib pairs (ASPs) with schizophrenia. Using 229 microsatellite markers at an average intermarker distance of 17.26 cM, followed in a second stage by a further 54 markers allowing the regions identified in stage 1 to be typed at an average spacing of 15 5.15 cM, Williams et al. considered results on chromosomes 4p, 18q, and Xcen as suggestive; however, given the scores, Williams et al. interpreted their results as suggesting that common genes of major effect (susceptibility ratio more than 3) are unlikely to exist for schizophrenia.

20 Similarly, Shaw et al., Am. J. Med. Genet. 81(5):364-76 (1998), in a genome-wide search for schizophrenia susceptibility genes, found that twelve chromosomes (1, 2, 4, 5, 8, 10, 11, 12, 13, 14, 16, and 22) had at least one region with a nominal P value <0.05, 25 that two of these chromosomes had a nominal P value <0.01 (chromosomes 13 and 16), and that five chromosomes (1, 2, 4, 11, and 13) had at least one marker with a lod score >2.0, suggesting the existence of multiple loci that contribute to schizophrenia susceptibility.

30 As yet another example, multiple genes are thought to predispose to epilepsy.

Epilepsy is characterized by recurrent, paroxysmal disorders of cerebral function (seizures); that is, by sudden, brief attacks of altered consciousness, 35 motor activity, sensory phenomena, or inappropriate

behavior. The risk of developing epilepsy is 1% in the period from birth to age 20, and 3% at age 75.

Epilepsy is caused by excessive discharge of cerebral neurons. Clinical manifestations depend on the 5 type and location of discharge. In partial seizures, for example, the excess neuronal discharge is contained within one region of the cerebral cortex. Simple partial seizures consist of motor, sensory, or psychomotor phenomena without loss of consciousness; the specific 10 phenomenon reflects the affected area of the brain. In generalized seizures, the discharge bilaterally and diffusely involves the entire cortex. Sometimes a focal lesion of one part of a hemisphere activates the entire cerebrum bilaterally so rapidly that it produces a 15 generalized tonic-clonic seizure before a focal sign appears.

Epilepsy is a family of disorders. Those that are idiopathic are believed to have multiple genetic contributions. For example, idiopathic generalized 20 epilepsy (IGE) is characterized by recurring generalized seizures in the absence of detectable brain lesions and/or metabolic abnormalities. Twin and family studies suggest that genetic factors play a key part in its etiology. Although a mutation in the CACNB4 gene can cause 25 the disorder, linkage to 8q24, Zara et al., Hum. Molec. Genet. 4: 1201-1207(1995), 3q26 and 14q23, Sander et al., Hum. Molec. Genet. 9:1465-1472 (2000), and 2q36 has been also demonstrated, with a multilocus model appearing to fit best the observed familial patterns.

30 Polygenic contributions to the etiology of various neurologic cancers have similarly been described.

For example, gliomas account for 45% of intracranial tumors, and multiple loci have been implicated in its development, with losses of chromosome 17p, increase 35 in copy number of chromosome 7, structural abnormalities of

chromosomes 9p and 19q, and genes on chromosome 10 among the suspects.

- Other significant diseases of brain and nervous tissue are also believed to have a genetic, typically
- 5 polygenic, etiologic component. These diseases include, for example, Parkinson's disease, dementia with Lewy bodies, frontotemporal dementia, corticobasal ganglionic degeneration, progressive supranuclear palsy, prion diseases (Creutzfeld-Jakob, Gerstmann-Straussler-Shenker,
- 10 familial fatal insomnia), Tourette's Syndrome, corticobasal degeneration, multiple system atrophy, striatonigral degeneration, Shy-Drager syndrome, olivopontocerebellar atrophy, spinocerebellar ataxia, Friedreich ataxia, ataxiatelangiectasia, amyotrophic lateral sclerosis, bulbospinal
- 15 atrophy (Kennedy's syndrome), spinal muscular atrophy, neuronal storage diseases (sphingolipid, mucopolysaccharide, mucolipid), leukodystrophy, Krabbe disease, metachromic leukodystrophy, adrenoleukodystrophy, Pelizaeus-Merzbacher disease, Canavan disease,
- 20 mitochondrial encephalomyopathy, Leigh disease, neurofibromatosis (Type I and Type II), tuberous sclerosis, paraneoplastic syndrome, subacute cerebellar degeneration, subacute sensory neuropathy, opsoclonus/myoclonus, retinal degeneration, stiff-man syndrome and Von Hippel-Lindau
- 25 disease.

- Many neurologic cancers other than gliomas have also been shown or suspected to have genetic bases or contributions. Among these cancers are astrocytoma, fibrillary astrocytoma, pilocytic astrocytoma,
- 30 pleomorphic xanthoastrocytoma, oligodendrogloma, ependymoma, gangliocytoma, ganglioglioma, medulloblastoma, primary brain germ cell tumor, pineocytoma, pineoblastoma, and meningioma.

- Other disorders of brain and central nervous system that likely have genetic components include the

- various forms of neural deafness, catatonia, depression, bipolar (manic-depressive) disorder, Wilson's Disease, Pick disease, neuromyelitis optica (Devic disease), central pontine myelinolysis, Marchiafava-Bignami disease,
- 5 Guillain-Barre syndrome, sleep disorders (insomnia, myoclonus, narcolepsy, cataplexy, sleep apnea), amnesia, aphasias (including Broca's aphasia and Wernicke's aphasia), cortical blindness, visual agnosia, auditory agnosia, and Kluver-Bucy syndrome.
- 10 The human genome-derived single exon nucleic acid probes and microarrays of the present invention are useful for predicting, diagnosing, grading, staging, monitoring and prognosing diseases of human brain, particularly those diseases with polygenic etiology. With each of the single
- 15 exon probes described herein shown to be expressed at detectable levels in human brain, and with about 2/3 of the probes identifying novel genes, the single exon microarrays of the present invention provide exceptionally high informational content for such studies.
- 20 For example, diagnosis (including differential diagnosis among clinically indistinguishable disorders), staging, and/or grading of a disease can be based upon the quantitative relatedness of a patient gene expression profile to one or more reference expression profiles known
- 25 to be characteristic of a given neurologic disease, or to specific grades or stages thereof.
- In one embodiment, the patient gene expression profile is generated by hybridizing nucleic acids obtained directly or indirectly from transcripts expressed in the
- 30 patient's brain (or other CNS tissues, including cultured tissues) to the genome-derived single exon microarray of the present invention. Reference profiles are be obtained similarly by hybridizing nucleic acids from individuals with known disease. Methods for quantitatively relating
- 35 gene expression profiles, without regard to the function of

the protein encoded by the gene, are disclosed in WO 99/58720, incorporated herein by reference in its entirety.

In another approach, the genome-derived single exon probes and microarrays of the present invention can be used to interrogate genomic DNA, rather than pools of expressed message; this latter approach permits predisposition to and/or prognosis of neurologic disease to be assessed through the massively parallel determination of altered copy number, deletion, or mutation in the patient's genome of exons known to be expressed in human brain. The algorithms set forth in WO 99/58720 can be applied to such genomic profiles without regard to the function of the protein encoded by the interrogated gene.

The utility is specific to the probe; at sufficiently high hybridization stringency, which stringencies are well known in the art - see Ausubel et al. and Maniatis et al. - each probe reports the level of expression of message specifically containing that ORF.

It should be appreciated, however, that the probes of the present invention, for which expression in the brain has been demonstrated are useful for both measurement in the brain and for survey of expression in other tissues.

Significant among such advantages is the presence of probes for novel genes.

As mentioned above and further detailed in Examples 1 and 2, the methods described enable ORFs which are not present in existing expression databases to be identified. And the fewer the number of tissues in which the ORF can be shown to be expressed, the more likely the ORF will prove to be part of a novel gene: as further discussed in Example 2, ORFs whose expression was measurable in only a single of the tested tissues were represented in existing expression databases at a rate of only 11%, whereas 36% of ORFs whose expression was

measurable in 9 tissues were present in existing expression databases, and fully 45% of those ORFs expressed in all ten tested tissues were present in existing expressed sequence databases.

5        Either as tools for measuring gene expression or tools for surveying gene expression, the genome-derived single exon probes of the present invention have significant advantages over the cDNA or EST-based probes that are currently available for achieving these utilities.

10      The genome-derived single exon probes of the present invention are useful in constructing genome-derived single exon microarrays; the genome-derived single exon microarrays, in turn, are useful devices for measuring and for surveying gene expression in the human.

15      Gene expression analysis using microarrays – conventionally using microarrays having probes derived from expressed message – is well-established as useful in the biological research arts (see Lockhart et al. *Nature* 405, 827-836).

20      Microarrays have been used to determine gene expression profiles in cells in response to drug treatment (see, for example, Kaminski et al., "Global Analysis of Gene Expression in Pulmonary Fibrosis Reveals Distinct Programs Regulating Lung Inflammation and Fibrosis," *Proc.*

25 *Natl. Acad. Sci. USA* 97(4):1778-83 (2000); Bartosiewicz et al., "Development of a Toxicological Gene Array and Quantitative Assessment of This Technology," *Arch. Biochem. Biophys.* 376(1):66-73 (2000)), viral infection (see for example, Geiss et al., "Large-scale Monitoring of Host Cell

30 Gene Expression During HIV-1 Infection Using cDNA Microarrays," *Virology* 266(1):8-16 (2000)) and during cell processes such as differentiation, senescence and apoptosis (see, for example, Shelton et al., "Microarray Analysis of Replicative Senescence," *Curr. Biol.* 9(17):939-45 (1999);

35 Voehringer et al., "Gene Microarray Identification of Redox

and Mitochondrial Elements That Control Resistance or Sensitivity to Apoptosis," *Proc. Natl. Acad. Sci. USA* 97(6):2680-5 (2000)).

- Microarrays have also been used to determine  
5 abnormal gene expression in diseased tissues (see, for example, Alon et al., "Broad Patterns of Gene Expression Revealed by Clustering Analysis of Tumor and Normal Colon Tissues Probed by Oligonucleotide Arrays," *Proc. Natl. Acad. Sci. USA* 96(12):6745-50 (1999); Perou et al.,  
10 "Distinctive Gene Expression Patterns in Human Mammary Epithelial Cells and Breast Cancers, *Proc. Natl. Acad. Sci. USA* 96(16):9212-7 (1999); Wang et al., "Identification of Genes Differentially Over-expressed in Lung Squamous Cell Carcinoma Using Combination of cDNA Subtraction and  
15 Microarray Analysis," *Oncogene* 19(12):1519-28 (2000); Whitney et al., "Analysis of Gene Expression in Multiple Sclerosis Lesions Using cDNA Microarrays," *Ann. Neurol.* 46(3):425-8 (1999)), in drug discovery screens (see, for example, Scherf et al., "A Gene Expression Database for the  
20 Molecular Pharmacology of Cancer," *Nat. Genet.* 24(3):236-44 (2000)) and in diagnosis to determine appropriate treatment strategies (see, for example, Sgroi et al., "In vivo Gene Expression Profile Analysis of Human Breast Cancer Progression," *Cancer Res.* 59(22):5656-61 (1999)).

- 25 In microarray-based gene expression screens of pharmacological drug candidates upon cells, each probe provides specific useful data. In particular, it should be appreciated that even those probes that show no change in expression are as informative as those that do change,  
30 serving, in essence, as negative controls.

- For example, where gene expression analysis is used to assess toxicity of chemical agents on cells, the failure of the agent to change a gene's expression level is evidence that the drug likely does not affect the pathway  
35 of which the gene's expressed protein is a part.

Analogously, where gene expression analysis is used to assess side effects of pharmacological agents – whether in lead compound discovery or in subsequent screening of lead compound derivatives – the inability of the agent to alter 5 a gene's expression level is evidence that the drug does not affect the pathway of which the gene's expressed protein is a part.

WO 99/58720 provides methods for quantifying the relatedness of a first and second gene expression profile 10 and for ordering the relatedness of a plurality of gene expression profiles. The methods so described permit useful information to be extracted from a greater percentage of the individual gene expression measurements from a microarray than methods previously used in the art.

15 Other uses of microarrays are described in Gerhold et al., *Trends Biochem. Sci.* 24(5):168-173 (1999) and Zweiger, *Trends Biotechnol.* 17(11):429-436 (1999); Schena et al.

The invention particularly provides genome- 20 derived single-exon probes known to be expressed in brain.

The individual single exon probes can be provided in the form of substantially isolated and purified nucleic acid, typically, but not necessarily, in a quantity sufficient to perform a hybridization reaction.

25 Such nucleic acid can be in any form directly hybridizable to the message that contains the probe's ORF, such as double stranded DNA, single-stranded DNA complementary to the message, single-stranded RNA complementary to the message, or chimeric DNA/RNA molecules 30 so hybridizable. The nucleic acid can alternatively or additionally include either nonnative nucleotides, alternative internucleotide linkages, or both, so long as complementary binding can be obtained. For example, probes can include phosphorothioates, methylphosphonates, 35 morpholino analogs, and peptide nucleic acids (PNA), as are

described, for example, in U.S. Patent Nos. 5,142,047; 5,235,033; 5,166,315; 5,217,866; 5,184,444; 5,861,250.

Usefully, however, such probes are provided in a form and quantity suitable for amplification, where the  
5 amplified product is thereafter to be used in the hybridization reactions that probe gene expression. Typically, such probes are provided in a form and quantity suitable for amplification by PCR or by other well known amplification technique. One such technique additional to  
10 PCR is rolling circle amplification, as is described, *inter alia*, in U.S. Patent Nos. 5,854,033 and 5,714,320 and international patent publications WO 97/19193 and WO 00/15779. As is well understood, where the probes are to be provided in a form suitable for amplification, the  
15 range of nucleic acid analogues and/or internucleotide linkages will be constrained by the requirements and nature of the amplification enzyme.

Where the probe is to be provided in form suitable for amplification, the quantity need not be  
20 sufficient for direct hybridization for gene expression analysis, and need be sufficient only to function as an amplification template, typically at least about 1, 10 or 100 pg or more.

Each discrete amplifiable probe can also be  
25 packaged with amplification primers, either in a single composition that comprises probe template and primers, or in a kit that comprises such primers separately packaged therefrom. As earlier mentioned, the ORF-specific 5' primers used for genomic amplification can have a first  
30 common sequence added thereto, and the ORF-specific 3' primers used for genomic amplification can have a second, different, common sequence added thereto, thus permitting, in this embodiment, the use of a single set of 5' and 3' primers to amplify any one of the probes. The probe  
35 composition and/or kit can also include buffers, enzyme,

etc., required to effect amplification.

- As mentioned earlier, when intended for use on a genome-derived single exon microarray of the present invention, the genome-derived single exon probes of the 5 present invention will typically average at least about 100, 200, 300, 400 or 500 bp in length, including (and typically, but not necessarily centered about) the ORF. Furthermore, when intended for use on a genome-derived single exon microarray of the present invention, the 10 genome-derived single exon probes of the present invention will typically not contain a detectable label.

When intended for use in solution phase hybridization, however – that is, for use in a hybridization reaction in which the probe is not first 15 bound to a support substrate (although the target may indeed be so bound) – length constraints that are imposed in microarray-based hybridization approaches will be relaxed, and such probes will typically be labeled.

- In such case, the only functional constraint that 20 dictates the minimum size of such probe is that each such probe must be capable of specifically identifying in a hybridization reaction the exon from which it is drawn. In theory, a probe of as little as 17 nucleotides is capable of uniquely identifying its cognate sequence in the human 25 genome. For hybridization to expressed message – a subset of target sequence that is much reduced in complexity as compared to genomic sequence – even fewer nucleotides are required for specificity.

Therefore, the probes of the present invention 30 can include as few as 20, 25 or 50 bp or ORF, or more. In particular embodiments, the ORF sequences are given in SEQ ID NOS. 12,822 - 25,434, respectively, for probe SEQ ID NOS. 1 - 12,821. The minimum amount of ORF required to be included in the probe of the present invention in order to 35 provide specific signal in either solution phase or

microarray-based hybridizations can readily be determined for each of ORF SEQ ID NOS. 12,822 - 25,434 individually by routine experimentation using standard high stringency conditions.

- 5        Such high stringency conditions are described, *inter alia*, in Ausubel et al. and Maniatis et al. For microarray-based hybridization, standard high stringency conditions can usefully be 50% formamide, 5X SSC, 0.2 µg/µl poly(dA), 0.2 µg/µl human c<sub>o</sub>t1 DNA, and 0.5 % SDS, in a  
10      humid oven at 42°C overnight, followed by successive washes of the microarray in 1X SSC, 0.2% SDS at 55°C for 5 minutes, and then 0.1X SSC, 0.2% SDS, at 55°C for 20 minutes. For solution phase hybridization, standard high stringency conditions can usefully be aqueous hybridization  
15      at 65°C in 6X SSC. Lower stringency conditions, suitable for cross-hybridization to mRNA encoding structurally- and functionally-related proteins, can usefully be the same as the high stringency conditions but with reduction in temperature for hybridization and washing to room  
20      temperature (approximately 25°C).

When intended for use in solution phase hybridization, the maximum size of the single exon probes of the present invention is dictated by the proximity of other expressed exons in genomic DNA: although each single  
25      exon probe can include intergenic and/or intronic material contiguous to the ORF in the human genome, each probe of the present invention will include portions of only one expressed exon.

Thus, each single exon probe will include no more  
30      than about 25 kb of contiguous genomic sequence, more typically no more than about 20 kb of contiguous genomic sequence, more usually no more than about 15 kb, even more usually no more than about 10 kb. Usually, probes that are maximally about 5 kb will be used, more typically no more  
35      than about 3 kb.

It will be appreciated that the Sequence Listing appended hereto presents, by convention, only that strand of the probe and ORF sequence that can be directly translated reading from 5' to 3' end. As would be well understood by one of skill in the art, single stranded probes must be complementary in sequence to the ORF as present in an mRNA; it is well within the skill in the art to determine such complementary sequence. It will further be understood that double stranded probes can be used in both solution-phase hybridization and microarray-based hybridization if suitably denatured.

Thus, it is an aspect of the present invention to provide single-stranded nucleic acid probes that have sequence complementary to those described herein above and below, and double-stranded probes one strand of which has sequence complementary to the probes described herein.

The probes can, but need not, contain intergenic and/or intronic material that flanks the ORF, on one or both sides, in the same linear relationship to the ORF that the intergenic and/or intronic material bears to the ORF in genomic DNA. The probes do not, however, contain nucleic acid derived from more than one expressed ORF.

And when intended for use in solution hybridization, the probes of the present invention can usefully have detectable labels. Nucleic acid labels are well known in the art, and include, *inter alia*, radioactive labels, such as  $^3\text{H}$ ,  $^{32}\text{P}$ ,  $^{33}\text{P}$ ,  $^{35}\text{S}$ ,  $^{125}\text{I}$ ,  $^{131}\text{I}$ ; fluorescent labels, such as Cy3, Cy5, Cy5.5, Cy7, SYBR®

Green and other labels described in Haugland, *Handbook of Fluorescent Probes and Research Chemicals*, 7th ed., Molecular Probes Inc., Eugene, OR (2000), or fluorescence resonance energy transfer tandem conjugates thereof; labels suitable for chemiluminescent and/or enhanced chemiluminescent detection; labels suitable for ESR and NMR detection; and labels that include one member

of a specific binding pair, such as biotin, digoxigenin, or the like.

The probes, either in quantity sufficient for hybridization or sufficient for amplification, can be  
5 provided in individual vials or containers.

Alternatively, such probes can usefully be packaged as a plurality of such individual genome-derived single exon probes.

When provided as a collection of plural  
10 individual probes, the probes are typically made available in amplifiable form in a spatially-addressable ordered set, typically one per well of a microtiter dish. Although a 96 well microtiter plate can be used, greater efficiency is obtained using higher density arrays.

15 If, as earlier mentioned, the ORF-specific 5' primers used for genomic amplification had a first common sequence added thereto, and the ORF-specific 3' primers used for genomic amplification had a second, different, common sequence added thereto, a single set of  
20 5' and 3' primers can be used to amplify all of the probes from the amplifiable ordered set.

Such collections of genome-derived single exon probes can usefully include a plurality of probes chosen for the common attribute of expression in the human brain.

25 In such defined subsets, typically at least 50, 60, 75, 80, 85, 90 or 95% or more of the probes will be chosen by their expression in the defined tissue or cell type.

The single exon probes of the present invention,  
30 as well as fragments of the single exon probes comprising selectively hybridizable portions of the probe ORF, can be used to obtain the full length cDNA that includes the ORF by (i) screening of cDNA libraries; (ii) rapid amplification of cDNA ends ("RACE"); or (iii) other  
35 conventional means, as are described, *inter alia*, in

Ausubel et al. and Maniatis et al.

- It is another aspect of the present invention to provide genome-derived single exon nucleic acid microarrays useful for gene expression analysis, where the term 5 "microarray" has the meaning given in the definitional section of this description, *supra*.

- The invention particularly provides genome-derived single-exon nucleic acid microarrays comprising a plurality of probes known to be expressed in human brain. 10 In preferred embodiments, the present invention provides human genome-derived single exon microarrays comprising a plurality of probes drawn from the group consisting of SEQ ID NOS.: 1 - 12,821.

- When used for gene expression analysis, the 15 genome-derived single exon microarrays provide greater physical informational density than do the genome-derived single exon microarrays that have lower percentages of probes known to be expressed commonly in the tested tissue. At a fixed probe density, for example, a given microarray 20 surface area of the defined subset genome-derived single exon microarray can yield a greater number of expression measurements. Alternatively, at a given probe density, the same number of expression measurements can be obtained from a smaller substrate surface area. Alternatively, at a 25 fixed probe density and fixed surface area, probes can be provided redundantly, providing greater reliability in signal measurement for any given probe. Furthermore, with a higher percentage of probes known to be expressed in the assayed tissue, the dynamic range of the detection means 30 can be adjusted to reveal finer levels discrimination among the levels of expression.

- Although particularly described with respect to their utility as probes of gene expression, particularly as probes to be included on a genome-derived single exon 35 microarray, each of the nucleic acids having SEQ ID NOS.: 1

- 12,821 contains an open-reading frame, set forth respectively in SEQ ID NOS.: 12,822 - 25,434, that encodes a protein domain. Thus, each of SEQ ID NOS. 1 - 12,821 can be used, or that portion thereof in SEQ ID NOS. 12,822 - 5 25,434 used, to express a protein domain by standard *in vitro* recombinant techniques. See Ausubel et al. and Maniatis et al.

Additionally, kits are available commercially that readily permit such nucleic acids to be expressed as 10 protein in bacterial cells, insect cells, or mammalian cells, as desired (e.g., HAT™ Protein Expression & Purification System, ClonTech Laboratories, Palo Alto, CA; Adeno-X™ Expression System, ClonTech Laboratories, Palo Alto, CA; Protein Fusion & Purification (pMAL™) System, New 15 England Biolabs, Beverly, MA)

Furthermore, shorter peptides can be chemically synthesized using commercial peptide synthesizing equipment and well known techniques. Procedures are described, *inter alia*, in Chan et al. (eds.), Fmoc Solid Phase Peptide Synthesis: A Practical Approach (Practical Approach Series, (Paper)), Oxford Univ. Press (March 2000) (ISBN: 0199637245); Jones, Amino Acid and Peptide Synthesis (Oxford Chemistry Primers, No 7), Oxford Univ. Press (August 1992) (ISBN: 0198556683); and Bodanszky, Principles of Peptide Synthesis (Springer Laboratory), Springer Verlag (December 1993) (ISBN: 0387564314).

It is, therefore, another aspect of the invention to provide peptides comprising an amino acid sequence translated from SEQ ID NOS.: 12,822 - 25,434. Such amino 30 acid sequences are set out in SEQ ID NOS: 25,435 - 37,811. Any such recombinantly-expressed or synthesized peptide of at least 8, and preferably at least about 15, amino acids, can be conjugated to a carrier protein and used to generate antibody that recognizes the peptide. Thus, it is a 35 further aspect of the invention to provide peptides that

have at least 8, preferably at least 15, consecutive amino acids.

The following examples are offered by way of  
5 illustration and not by way of limitation.

EXAMPLE 1

Preparation of Single Exon Microarrays from ORFs Predicted  
in Human Genomic Sequence

10

Bioinformatics Results

All human BAC sequences in fewer than 10 pieces that had been accessioned in a five month period immediately preceding this study were downloaded from 15 GenBank. This corresponds to ~2200 clones, totaling ~350 MB of sequence, or approximately 10% of the human genome.

After masking repetitive elements using the program CROSS\_MATCH, the sequence was analyzed for open reading frames using three separate gene finding programs. 20 The three programs predict genes using independent algorithmic methods developed on independent training sets: GRAIL uses a neural network, GENEFINDER uses a hidden Markoff model, and DICTION, a program proprietary to Genetics Institute, operates according to a different heuristic. The results of all three programs were used to 25 create a prediction matrix across the segment of genomic DNA.

The three gene finding programs yielded a range of results. GRAIL identified the greatest percentage of 30 genomic sequence as putative coding region, 2% of the data analyzed. GENEFINDER was second, calling 1%, and DICTION yielded the least putative coding region, with 0.8% of genomic sequence called as coding region.

The consensus data were as follows. GRAIL and 35 GENEFINDER agreed on 0.7% of genomic sequence, GRAIL and

DICTION agreed on 0.5% of genomic sequence, and the three programs together agreed on 0.25% of the data analyzed. That is, 0.25% of the genomic sequence was identified by all three of the programs as containing putative coding 5 region.

ORFs predicted by any two of the three programs ("consensus ORFs") were assorted into "gene bins" using two criteria: (1) any 7 consecutive exons within a 25 kb window were placed together in a bin as likely contributing to a 10 single gene, and (2) all ORFs within a 25 kb window were placed together in a bin as likely contributing to a single gene if fewer than 7 exons were found within the 25 kb window.

15 PCR

The largest ORF from each gene bin that did not span repetitive sequence was then chosen for amplification, as were all consensus ORFs longer than 500 bp. This method approximated one exon per gene; however, a number of genes 20 were found to be represented by multiple elements.

Previously, we had determined that DNA fragments fewer than 250 bp in length do not bind well to the amino-modified glass surface of the slides used as support substrate for construction of microarrays; therefore, 25 amplicons were designed in the present experiments to approximate 500 bp in length.

Accordingly, after selecting the largest ORF per gene bin, a 500 bp fragment of sequence centered on the ORF was passed to the primer picking software, PRIMER3 30 (available online for use at <http://www-genome.wi.mit.edu/cgi-bin/primer/>). A first additional sequence was commonly added to each ORF-unique 5' primer, and a second, different, additional sequence was commonly added to each ORF-unique 3' primer, to permit 35 subsequent reamplification of the amplicon using a single

set of "universal" 5' and 3' primers, thus immortalizing the amplicon. The addition of universal priming sequences also facilitates sequence verification, and can be used to add a cloning site should some ORFs be found to warrant  
5 further study.

The ORFs were then PCR amplified from genomic DNA, verified on agarose gels, and sequenced using the universal primers to validate the identity of the amplicon to be spotted in the microarray.

10 Primers were supplied by Operon Technologies (Alameda, CA). PCR amplification was performed by standard techniques using human genomic DNA (Clontech, Palo Alto, CA) as template. Each PCR product was verified by SYBR® green (Molecular Probes, Inc., Eugene, OR) staining of  
15 agarose gels, with subsequent imaging by Fluorimager (Molecular Dynamics, Inc., Sunnyvale, CA). PCR amplification was classified as successful if a single band appeared.

20 The success rate for amplifying ORFs of interest directly from genomic DNA using PCR was approximately 75%. FIG. 5 graphs the distribution of predicted ORF (exon) length and distribution of amplified PCR products, with ORF length shown in red and PCR product length shown in blue (which may appear black in the figure). Although the range  
25 of ORF sizes is readily seen to extend to beyond 900 bp, the mean predicted exon size was only 229 bp, with a median size of 150 bp (n=9498). With an average amplicon size of 475 ± 25 bp, approximately 50% of the average PCR amplification product contained predicted coding region,  
30 with the remaining 50% of the amplicon containing either intron, intergenic sequence, or both.

Using a strategy predicated on amplifying about 500 bp, it was found that long exons had a higher PCR failure rate. To address this, the bioinformatics process  
35 was adjusted to amplify 1000, 1500 or 2000 bp fragments

from exons larger than 500 bp. This improved the rate of successful amplification of exons exceeding 500 bp, constituting about 9.2% of the exons predicted by the gene finding algorithms.

- 5        Approximately 75% of the probes disposed on the array (90% of those that successfully PCR amplified) were sequence-verified by sequencing in both the forward and reverse direction using MegaBACE sequencer (Molecular Dynamics, Inc., Sunnyvale, CA), universal primers, and  
10 standard protocols.

Some genomic clones (BACs) yielded very poor PCR and sequencing results. The reasons for this are unclear, but may be related to the quality of early draft sequence or the inclusion of vector and host contamination in some  
15 submitted sequence data.

- Although the intronic and intergenic material flanking coding regions could theoretically interfere with hybridization during microarray experiments, subsequent empirical results demonstrated that differential expression  
20 ratios were not significantly affected by the presence of noncoding sequence. The variation in exon size was similarly found not to affect differential expression ratios significantly; however, variation in exon size was observed to affect the absolute signal intensity (data not  
25 shown).

- The 350 MB of genomic DNA was, by the above-described process, reduced to 9750 discrete probes, which were spotted in duplicate onto glass slides using commercially available instrumentation (MicroArray GenII  
30 Spotter and/or MicroArray GenIII Spotter, Molecular Dynamics, Inc., Sunnyvale, CA). Each slide additionally included either 16 or 32 *E. coli* genes, the average hybridization signal of which was used as a measure of background biological noise.

- 35        Each of the probe sequences was BLASTed against

the human EST data set, the NR data set, and SwissProt GenBank (May 7, 1999 release 2.0.9).

One third of the probe sequences (as amplified) produced an exact match (BLAST Expect ("E") values less than  $1 e^{-100}$ ) to either an EST (20% of sequences) or a known mRNA (13% of sequences). A further 22% of the probe sequences showed some homology to a known EST or mRNA (BLAST E values from  $1 e^{-5}$  to  $1 e^{-99}$ ). The remaining 45% of the probe sequences showed no significant sequence homology to any expressed, or potentially expressed, sequences present in public databases.

All of the probe sequences (as amplified) were then analyzed for protein similarities with the SwissProt database using BLASTX, Gish et al., *Nature Genet.* 3:266 (1993). The predicted functional breakdowns of the 2/3 of probes identical or homologous to known sequences are presented in Table 1.

Table 1

Function of Predicted ORFs As Deduced From Comparative Sequence Analysis			
Total	V6 chip	V7 chip	Function Predicted from Comparative Sequence Analysis
211	96	115	Receptor
120	43	77	Zinc Finger
30	11	19	Homeobox
25	9	16	Transcription Factor
17	11	7	Transcription
118	57	61	Structural
95	39	56	Kinase
36	18	18	Phosphatase
83	31	52	Ribosomal

45	19	26	Transport
21	17	14	Growth Factor
17	12	5	Cytochrome
50	33	17	Channel

As can be seen, the two most common types of genes were transcription factors and receptors, making up 2.2% and 1.8% of the arrayed elements, respectively.

5

#### EXAMPLE 2

#### Gene Expression Measurements From Genome-Derived Single Exon Microarrays

10

The two genome-derived single exon microarrays prepared according to Example 1 were hybridized in a series of simultaneous two-color fluorescence experiments to (1) 15 Cy3-labeled cDNA synthesized from message drawn individually from each of brain, heart, liver, fetal liver, placenta, lung, bone marrow, HeLa, BT 474, or HBL 100 cells, and (2) Cy5-labeled cDNA prepared from message pooled from all ten tissues and cell types, as a control in 20 each of the measurements. Hybridization and scanning were carried out using standard protocols and Molecular Dynamics equipment.

Briefly, mRNA samples were bought from commercial sources (Clontech, Palo Alto, CA and Amersham Pharmacia Biotech (APB)). Cy3-dCTP and Cy5-dCTP (both from APB) were incorporated during separate reverse transcriptions of 1 µg of polyA<sup>+</sup> mRNA performed using 1 µg oligo(dT)12-18 primer and 2 µg random 9mer primers as follows. After heating to 70°C, the RNA:primer mixture was snap cooled on ice. After 30 snap cooling on ice, added to the RNA to the stated final concentration was: 1X Superscript II buffer, 0.01 M DTT,

100 μM dATP, 100 μM dGTP, 100 μM dTTP, 50 μM dCTP, 50 μM Cy3-dCTP or Cy5-dCTP 50 μM, and 200 U Superscript II enzyme. The reaction was incubated for 2 hours at 42°C. After 2 hours, the first strand cDNA was isolated by adding 5 1 U Ribonuclease H, and incubating for 30 minutes at 37°C. The reaction was then purified using a Qiagen PCR cleanup column, increasing the number of ethanol washes to 5. Probe was eluted using 10 mM Tris pH 8.5.

Using a spectrophotometer, probes were measured 10 for dye incorporation. Volumes of both Cy3 and Cy5 cDNA corresponding to 50 pmoles of each dye were then dried in a Speedvac, resuspended in 30 μl hybridization solution containing 50% formamide, 5X SSC, 0.2 μg/μl poly(dA), 0.2 μg/μl human c<sub>o</sub>t1 DNA, and 0.5 % SDS.

15 Hybridizations were carried out under a coverslip, with the array placed in a humid oven at 42°C overnight. Before scanning, slides were washed in 1X SSC, 0.2% SDS at 55°C for 5 minutes, followed by 0.1X SSC, 0.2% SDS, at 55°C for 20 minutes. Slides were briefly dipped in 20 water and dried thoroughly under a gentle stream of nitrogen.

Slides were scanned using a Molecular Dynamics Gen3 scanner, as described. Schena (ed.), Microarray Biochip: Tools and Technology, Eaton Publishing 25 Company/BioTechniques Books Division (2000) (ISBN: 1881299376).

Although the use of pooled cDNA as a reference permitted the survey of a large number of tissues, it attenuates the measurement of relative gene expression, 30 since every highly expressed gene in the tissue/cell type-specific fluorescence channel will be present to a level of at least 10% in the control channel. Because of this fact, both signal and expression ratios (the latter hereinafter, "expression" or "relative expression") for each probe were 35 normalized using the average ratio or average signal,

respectively, as measured across the whole slide.

Data were accepted for further analysis only when signal was at least three times greater than biological noise, the latter defined by the average signal produced by 5 the *E. coli* control genes.

The relative expression signal for these probes was then plotted as function of tissue or cell type, and is presented in FIG. 6.

FIG. 6 shows the distribution of expression 10 across a panel of ten tissues. The graph shows the number of sequence-verified products that were either not expressed ("0"), expressed in one or more but not all tested tissues ("1" - "9"), and expressed in all tissues tested ("10").

15 Of 9999 arrayed elements on the two microarrays (including positive and negative controls and "failed" products), 2353 (51%) were expressed in at least one tissue or cell type. Of the gene elements showing significant signal - where expression was scored as "significant" if 20 the normalized Cy3 signal was greater than 1, representing signal 5-fold over biological noise (0.2) - 39% (991) were expressed in all 10 tissues. The next most common class (15%) consisted of gene elements expressed in only a single tissue.

25 The genes expressed in a single tissue were further analyzed, and the results of the analyses are compiled in FIG. 7.

FIG. 7A is a matrix presenting the expression of all verified sequences that showed expression greater than 30 3 in at least one tissue. Each clone is represented by a column in the matrix. Each of the 10 tissues assayed is represented by a separate row in the matrix, and relative expression of a clone in that tissue is indicated at the respective node by intensity of green shading, with the 35 intensity legend shown in panel B. The top row of the

matrix ("EST Hit") contains "bioinformatic" rather than "physical" expression data - that is, presents the results returned by query of EST, NR and SwissProt databases using the probe sequence. The legend for "bioinformatic expression" (i.e., degree of homology returned) is presented in panel C. Briefly, white is known, black is novel, with gray depicting nonidentical with significant homology (white: E values < 1e-100; gray: E values from 1e-05 to 1e-99; black: E values > 1e-05).

As FIG. 7 readily shows, heart and brain were demonstrated to have the greatest numbers of genes that were shown to be uniquely expressed in the respective tissue. In brain, 200 uniquely expressed genes were identified; in heart, 150. The remaining tissues gave the following figures for uniquely expressed genes: liver, 100; lung, 70; fetal liver, 150; bone marrow, 75; placenta, 100; HeLa, 50; HBL, 100; and BT474, 50.

It was further observed that there were many more "novel" genes among those that were up-regulated in only one tissue, as compared with those that were down-regulated in only one tissue. In fact, it was found that ORFs whose expression was measurable in only a single of the tested tissues were represented in sequencing databases at a rate of only 11%, whereas 36% of the ORFs whose expression was measurable in 9 of the tissues were present in public databases. As for those ORFs expressed in all ten tissues, fully 45% were present in existing expressed sequence databases. These results are not unexpected, since genes expressed in a greater number of tissues have a higher likelihood of being, and thus of having been, discovered by EST approaches.

#### Comparison of Signal from Known and Unknown Genes

The normalized signal of the genes found to have high homology to genes present in the GenBank human EST

database were compared to the normalized signal of those genes not found in the GenBank human EST database. The data are shown in FIG. 8.

FIG. 8 shows the normalized Cy3 signal intensity 5 for all sequence-verified products with a BLAST Expect ("E") value of greater than 1e-30 (designated "unknown") upon query of existing EST, NR and SwissProt databases, and shows in blue the normalized Cy3 signal intensity for all sequence-verified products with a BLAST Expect value of 10 less than 1e-30 ("known"). Note that biological background noise has an averaged normalized Cy3 signal intensity of 0.2.

As expected, the most highly expressed of the ORFs were "known" genes. This is not surprising, since 15 very high signal intensity correlates with very commonly-expressed genes, which have a higher likelihood of being found by EST sequence.

However, a significant point is that a large number of even the high expressers were "unknown". Since 20 the genomic approach used to identify genes and to confirm their expression does not bias exons toward either the 3' or 5' end of a gene, many of these high expression genes will not have been detected in an end-sequenced cDNA library.

25 The significant point is that presence of the gene in an EST database is not a prerequisite for incorporation into a genome-derived microarray, and further, that arraying such "unknown" exons can help to assign function to as-yet undiscovered genes.

30

#### Verification of Gene Expression

To ascertain the validity of the approach described above to identify genes from raw genomic sequence, expression of two of the probes was assayed using 35 reverse transcriptase polymerase chain reaction (RT PCR)

and northern blot analysis.

Two microarray probes were selected on the basis of exon size, prior sequencing success, and tissue-specific gene expression patterns as measured by the microarray 5 experiments. The primers originally used to amplify the two respective ORFs from genomic DNA were used in RT PCR against a panel of tissue-specific cDNAs (Rapid-Scan gene expression panel 24 human cDNAs) (Origene Technologies, Inc., Rockville, MD).

10 Sequence AL079300\_1 was shown by microarray hybridization to be present in cardiac tissue, and sequence AL031734\_1 was shown by microarray experiment to be present in placental tissue (data not shown). RT-PCR on these two sequences confirmed the tissue-specific gene expression as 15 measured by microarrays, as ascertained by the presence of a correctly sized PCR product from the respective tissue type cDNAs.

Clearly, all microarray results cannot, and indeed should not, be confirmed by independent assay 20 methods, or the high throughput, highly parallel advantages of microarray hybridization assays will be lost. However, in addition to the two RT-PCR results presented above, the observation that 1/3 of the arrayed genes exist in expression databases provides powerful confirmation of the 25 power of our methodology – which combines bioinformatic prediction with expression confirmation using genome-derived single exon microarrays – to identify novel genes from raw genomic data.

To verify that the approach further provides 30 correct characterization of the expression patterns of the identified genes, a detailed analysis was performed of the microarrayed sequences that showed high signal in brain.

For this latter analysis, sequences that showed 35 high (normalized) signal in brain, but which showed very low (normalized) signal (less than 0.5, determined to be

biological noise) in all other tissues, were further studied. There were 82 sequences that fit these criteria, approximately 2% of the arrayed elements. The 10 sequences showing the highest signal in brain in microarray 5 hybridizations are detailed in Table 2, along with assigned function, if known or reasonably predicted.

Table 2

Function of the Most Highly Expressed Genes Expressed Only in Brain				
Microarray Sequence Name	Normal ized Signal	Expressi on Ratio	Homology to EST present in GenBank	Gene Function as described by GenBank
AP000217-1	5.2	+7.7	High	S-100 protein, b-chain, Ca <sup>2+</sup> binding protein expressed in central nervous system
AP000047-1	2.3		High	Unknown Function
AC006548-9	1.7		High	Similar to mouse membrane glyco-protein M6, expressed in central nervous system

AC007245-5	1.5		High	Similar to amphiphysin, a synaptic vesicle-associated protein. Ref 21
L44140-4	1.2	+2.0	High	Endothelial actin-binding protein found in nonmuscle filamin
AC004689-9	1.2	+3.5	High	Protein Phosphatase PP2A, neuronal/ downregulates activated protein kinases
AL031657-1	1.2	+3.0	High	Unknown function/ Contains the anhyrin motif, a common protein sequence motif
AC009266-2	1.1	+3.7	Low	Low homology to the Synaptotagmin I protein in rat/present at low levels throughout rat brain
AP000086-1	1.0	+2.7	Low	Unknown, very poor homology to collagen

AC004689-3	1.0		High	Protein Phosphatase PP2A, neuronal/ downregulates activated protein kinases
------------	-----	--	------	--

Of the ten sequences studied by these latter confirmatory approaches, eight were previously known. Of these eight, six had previously been reported to be 5 important in the central nervous system or brain. The exon giving the highest signal (AP00217-1) was found to be the gene encoding an S100B  $\text{Ca}^{2+}$  binding protein, reported in the literature to be highly and uniquely expressed in the central nervous system. Heizmann, *Neurochem. Res.* 9:1097 10 (1997).

A number of the brain-specific probe sequences (including AC006548-9, AC009266-2) did not have homology to any known human cDNAs in GenBank but did show homology to rat and mouse cDNAs. Sequences AC004689-9 and AC004689-3 15 were both found to be phosphatases present in neurons (Millward et al., *Trends Biochem. Sci.* 24(5):186-191 (1999)). Two microarray sequences, AP000047-1 and AP000086-1 have unknown function, with AP000086-1 being absent from GenBank. Functionality can now be narrowed 20 down to a role in the central nervous system for both of these genes, showing the power of designing microarrays in this fashion.

Next, the function of the chip sequences with the highest (normalized) signal intensity in brain, regardless 25 of expression in other tissues, was assessed. In this latter analysis, we found expression of many more common genes, since the sequences were not limited to those expressed only in brain. For example, looking at the 20 highest signal intensity spots in brain, 4 were similar to

tubulin (AC00807905; AF146191-2; AC007664-4; AF14191-2), 2 were similar to actin (AL035701-2; AL034402-1), and 6 were found to be homologous to glyceraldehyde-3-phosphate dehydrogenase (GAPDH) (AL035604-1; Z86090-1; AC006064-L, 5 AC006064-K; AC035604-3; AC006064-L). These genes are often used as controls or housekeeping genes in microarray experiments of all types.

Other interesting genes highly expressed in brain were a ferritin heavy chain protein, which is reported in 10 the literature to be found in brain and liver (Joshi et al., *J. Neurol. Sci.* 134(Suppl):52-56 (1995)), a result duplicated with the array. Other highly expressed chip sequences included a translation elongation factor 10 (AC007564-4), a DEAD-box homolog (AL023804-4), and a Y- 15 chromosome RNA-binding motif (Chai et al., *Genomics* 49(2):283-89 (1998)) (AC007320-3). A low homology analog (AP00123-1/2) to a gene, DSCR1, thought to be involved in trisomy 21 (Down's syndrome), showed high expression in both brain and heart, in agreement with the literature 20 (Fuentes et al., *Mol. Genet.* 4(10):1935-44 (1995)).

As a further validation of the approach, we selected the BAC AC006064 to be included on the array. This BAC was known to contain the GAPDH gene, and thus could be used as a control for the ORF selection process. 25 The gene finding and exon selection algorithms resulted in choosing 25 exons from BAC AC006064 for spotting onto the array, of which four were drawn from the GAPDH gene. Table 3 shows the comparison of the average expression ratio for the 4 exons from BAC006064 compared with the average 30 expression ratio for 5 different dilutions of a commercially available GAPDH cDNA (Clontech).

Table 3

Comparison of Expression Ratio, for each tissue, of GAPDH

	AC006064 (n = 4)	Control (n = 5)
Bone Marrow	-1.81 ± 0.11	-1.85 ± 0.08
Brain	-1.41 ± 0.11	-1.17 ± 0.05
BT474	1.85 ± 0.09	1.66 ± 0.12
Fetal Liver	-1.62 ± 0.07	-1.41 ± 0.05
HBL100	1.32 ± 0.05	2.64 ± 0.12
Heart	1.16 ± 0.09	1.56 ± 0.10
HeLa	1.11 ± 0.06	1.30 ± 0.15
Liver	-1.62 ± 0.22	-2.07 ±
Lung	-4.95 ± 0.93	-3.75 ± 0.21
Placenta	-3.56 ± 0.25	-3.52 ± 0.43

Each tissue shows excellent agreement between the experimentally chosen exons and the control, again demonstrating the validity of the present exon mining approach. In addition, the data also show the variability of expression of GAPDH within tissues, calling into question its classification as a housekeeping gene and utility as a housekeeping control in microarray experiments.

### EXAMPLE 3

Representation of Sequence and Expression Data as a "Mondrian"

15

For each genomic clone processed for microarray as above-described, a plethora of information was accumulated, including full clone sequence, probe sequence within the clone, results of each of the three gene finding programs, EST information associated with the probe

sequences, and microarray signal and expression for multiple tissues, challenging our ability to display the information.

Accordingly, we devised a new tool for visual  
5 display of the sequence with its attendant annotation which, in deference to its visual similarity to the paintings of Piet Mondrian, is hereinafter termed a "Mondrian". FIGS. 3 and 4 present the key to the information presented on a Mondrian.

10 FIG. 9 presents a Mondrian of BAC AC008172 (bases 25,000 to 130,000 shown), containing the carbamyl phosphate synthetase gene (AF154830.1). Purple background within the region shown as field 81 in FIG. 3 indicates all 37 known exons for this gene.

15 As can be seen, GRAIL II successfully identified 27 of the known exons (73%), GENEFINDER successfully identified 37 of the known exons (100%), while DICTION identified 7 of the known exons (19%).

20 Seven of the predicted exons were selected for physical assay, of which 5 successfully amplified by PCR and were sequenced. These five exons were all found to be from the same gene, the carbamyl phosphate synthetase gene (AF154830.1).

25 The five exons were arrayed, and gene expression measured across 10 tissues. As is readily seen in the Mondrian, the five chip sequences on the array show identical expression patterns, elegantly demonstrating the reproducibility of the system.

FIG. 10 is a Mondrian of BAC AL049839. We  
30 selected 12 exons from this BAC, of which 10 successfully sequenced, which were found to form between 5 and 6 genes. Interestingly, 4 of the genes on this BAC are protease inhibitors. Again, these data elegantly show that exons selected from the same gene show the same expression  
35 patterns, depicted below the red line. From this figure,

it is clear that our ability to find known genes is very good. A novel gene is also found from 86.6 kb to 88.6 kb, upon which all the exon finding programs agree. We are confident we have two exons from a single gene since they  
5 show the same expression patterns and the exons are proximal to each other. Backgrounds in the following colors indicate a known gene (top to bottom):  
red = kallistatin protease inhibitor (P29622);  
purple = plasma serine protease inhibitor (P05154);  
10 turquoise =  $\alpha 1$  anti-chymotrypsin (P01011); mauve = 40S ribosomal protein (P08865). Note that chip sequence 8 and 12 did not sequence verify.

15 EXAMPLE 4

Genome-Derived Single Exon Probes Useful For Measuring Human Gene Expression

The protocols set forth in Examples 1 and 2,  
20 supra, were applied to additional human genomic sequence as it became newly available in GenBank to identify unique exons in the human genome that could be shown to be expressed at significant levels in brain tissue.

These unique exons are within longer probe  
25 sequences. Each probe was completely sequenced on both strands prior to its use on a genome-derived single exon microarray; sequencing confirms the exact chemical structure of each probe. An added benefit of sequencing is that it placed us in possession of a set of single base-  
30 incremented fragments of the sequenced nucleic acid, starting from the sequencing primer 3' OH. (Since the single exon probes were first obtained by PCR amplification from genomic DNA, we were of course additionally in possession of an even larger set of single base incremented  
35 fragments of each of the 12,821 single exon probes, each

fragment corresponding to an extension product from one of the two amplification primers.)

The structures of the 12,821 unique single exon probes are clearly presented in the Sequence Listing as SEQ ID Nos.: 1 - 12,821. The 16 nt 5' primer sequence and 16 nt 3' primer sequence present on the amplicon are not included in the sequence listing. The sequences of the exons present within each of these probes is presented in the Sequence Listing as SEQ ID NOs.: 12,822 - 25,434, respectively. It will be noted that some amplicons have more than one exon, some exons are contained in more than one amplicon.

As detailed in Example 2, expression was demonstrated by disposing the amplicons as single exon probes on nucleic acid microarrays and then performing two-color fluorescent hybridization analysis; significant expression is based on a statistical confidence that the signal is significantly greater than negative biological control spots. The negative biological control is formed from spotted DNA sequences from a different species. Here, 32 sequences from E.Coli were spotted in duplicate to give a total of 64 spots.

For each hybridisation (each slide, each colour) the median value of the signal from all of the spots is determined. The normalised signal value is the arithmetic mean of the signal from duplicate spots divided by the population median.

Control spots are eliminated if there is more than a five-fold difference between each one of the duplicate spots raw signals.

The median of the signal from the remaining control spots is calculated and all subsequent calculations are done with normalised signals.

Control spots having a signal of greater than median + 2.4 (the value 2.4 is roughly 12 times the

observed standard deviation of control spot populations) are eliminated. Spots with such high signals are considered to be "outliers".

5 The mean and standard deviation of the modified control spot populations are calculated.

The mean + 3x the standard deviation (mean + (3\*SD)) is used as the signal threshold qualifier for that particular hybridisation. Thus, individual thresholds are determined for each channel and each hybridisation.

10 This means that, assuming that the data is distributed normally, there is a 99% confidence that any signal exceeding the threshold is significant.

The probes and their expression data are presented in Table 4, set forth respectively in Example 5.

15 Example 5 presents the subset of probes that is significantly expressed in the human heart and thus presents the subset of probes that was recognized to be useful for measuring expression of their cognate genes in human brain tissue.

20 The sequence of each of the exon probes identified by SEQ ID NOS.: 12,822 - 25,434 was individually used as a BLAST (or, for SWISSPROT, BLASTX) query to identify the most similar sequence in each of dbEST, SwissProt (BLASTX), and NR divisions of GenBank. Because 25 the query sequences are themselves derived from genomic sequence in GenBank, only nongenomic hits from NR were scored.

The smallest in value of the BLAST (or BLASTX) expect ("E") scores for each query sequence across the 30 three database divisions was used as a measure of the "expression novelty" of the probe's ORF. Table 4 is sorted in descending order based on this measure, reported as "Most Similar (top) Hit BLAST E Value". Those sequences for which no "Hit E Value" is listed are those exons which were 35 found to have no similar sequences.

As sorted, Table 4 thus lists its respective probes (by "AMPLICON SEQ ID NO.:" and additionally by the SEQ ID NO.: of the exon contained within the probe: "EXON SEQ ID NO.:") from least similar to sequences known to be expressed (i.e., highest BLAST E value), at the beginning of the table, to most similar to sequences known to be expressed (i.e., lowest BLAST E value), at the bottom of the table.

Table 4 further provides, for each listed probe, the accession number of the database sequence that yielded the "Most Similar (top) Hit BLAST E Value", along with the name of the database in which the database sequence is found ("Top Hit Database Source").

Table 4 further provides SEQ ID NOS.

corresponding to the predicted amino acid sequences where they have been determined for the probe and exon nucleotide sequences. These are set out as PEPTIDE SEQ ID NOS.: . The peptide sequences for a given exon are predicted as follows: Since each chip exon is a consensus sequence drawn from predictions from various exon finding programs (i.e. Grail, GeneFinder and GenScan), the multiple initial ORFs are first determined in a uniform way according to each prediction. In particular, the reading frame for predicting the first amino acid in the peptide sequence always starts with the first base of any codon and ends with the last base of non-termination codon. Next, for each strand of the exon, initial ORFs are merged into one or more final ORFs in an exhaustive process based on the following criteria: 1) the merging ORFs must be overlapping, and 2) the merging ORFs must be in the same frame.

The Sequence Listing, which is a superset of all of the data presented in Table 4, further includes, for each probe, the most similar hit, with accession number and BLAST E value, from each of the three queried databases.

Table 4 further lists, for each probe, a portion of the descriptor for the top hit ("Top Hit Descriptor") as provided in the sequence database. For those ORFs that are similar in sequence, but nonidentical to known sequences 5 (e.g., those with BLAST E values between about 1e-05 and 1e-100), the descriptor reveals the likely function of the protein encoded by the probe's ORF.

Using BLAST E value cutoffs of 1e-05 (i.e.,  $1 \times 10^{-5}$ ) and 1e-100 (i.e.,  $1 \times 10^{-100}$ ) as evidence of similarity 10 to sequences known to be expressed is of course arbitrary: in Example 2, *supra*, a BLAST E value of 1e-30 was used as the boundary when only two classes were to be defined for analysis (unknown,  $>1e-30$ ; known  $<1e-30$ ) (see also FIG. 8). Furthermore, even when the "Most Similar (Top) Hit BLAST E 15 Value" is low, e.g., less than about 1e-100 – which is probative evidence that the query sequence has previously been shown to be expressed – the top hit is highly unlikely exactly to match the probe sequence.

First, such expression entries typically will not 20 have the intronic and/or intergenic sequence present within the single exon probes listed in the Table. Second, even the ORF itself is unlikely in such cases to be present identically in the databases, since most of the EST and mRNA clones in existing databases include multiple exons, 25 without any indication of the location of exon boundaries.

As noted, the data presented in Table 4 represent a proper subset of the data present within the attached sequence listing. For each amplicon probe (SEQ ID NOS.: 1 - 12,821) and probe exon (SEQ ID NOS.: 12,822 - 25,434, 30 respectively), the sequence listing further provides, through iterated annotation fields <220> and <223>:

(a) the accession number of the BAC from which the sequence was derived ("MAP TO"), thus providing a link to the chromosomal map location and other information about 35 the genomic milieu of the probe sequence;

(b) the most similar sequence provided by BLAST query of the EST database, with accession number and BLAST E value for the "hit";

5 (c) the most similar sequence provided by BLAST query of the GenBank NR database, with accession number and BLAST E value for the "hit"; and

(d) the most similar sequence provided by BLASTX query of the SWISSPROT database, with accession number and BLAST E value for the "hit".

10

EXAMPLE 5

Genome-Derived Single Exon Probes Useful For Measuring Expression of Genes in Human Brain

15

Table 4 (536 pages) presents expression, homology, and functional information for the genome-derived single exon probes that are expressed significantly in human brain.

20

## CLAIMS

1. A spatially-addressable set of single exon nucleic acid probes for measuring gene expression in a sample derived from human brain comprising a plurality single exon nucleic probes, said probes comprising any one of the nucleotide sequences set out in SEQ ID NOs: 1 - 12,821 or a complementary sequence, or a portion of such a sequence.
- 10 2. A spatially-addressable set of single exon nucleic acid probes as claimed in claim 1 wherein each of said plurality of probes is separately and addressably amplifiable.
- 15 3. A spatially-addressable set of single exon nucleic acid probes as claimed in claim 1 wherein each of said plurality of probes is separately and addressably isolatable from said plurality.
- 20 4. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 3 wherein said probes comprise any one of the nucleotide sequences set out in SEQ ID NOS.: 12,822 - 25,434.
- 25 5. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 4, wherein each of said plurality of probes is amplifiable using at least one common primer.
- 30 6. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 5 wherein the set comprises between 50 - 20,000 single exon nucleic acid probes.
- 35 7. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 6, wherein the

average length of the single exon nucleic acid probes is between 200 and 500 bp.

8. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 7, wherein at least 5% of said single exon nucleic acid probes lack prokaryotic and bacteriophage vector sequence.
9. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 to 8, wherein at least 5% of said single exon nucleic acid probes lack homopolymeric stretches of A or T.
10. A spatially-addressable set of single exon nucleic acid probes as claimed in any of claims 1 - 9 characterised in that said set of probes is addressably disposed upon a substrate.
11. A spatially-addressable set of single exon nucleic acid probes as claimed in claim 10 wherein said substrate is selected from glass, amorphous silicon, crystalline silicon and plastic.
12. A microarray comprising a spatially addressable set of single exon nucleic acid probes as claimed in any of claims 1 - 11.
13. A single exon nucleic acid probe for measuring human gene expression in a sample derived from human brain comprising a nucleotide sequence as set out in any of SEQ ID NOs.: 1 - 12,821 or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid molecule expressed in the human brain.

14. A single exon nucleic acid probe as claimed in claim 13 comprising a nucleotide sequence as set out in any of SEQ ID NOs.: 12,822 - 25,434 or a complementary sequence or a fragment thereof.

5

15. A single exon nucleic acid probe for measuring human gene expression in a sample derived from human brain which is a nucleic acid molecule having a sequence encoding a peptide comprising a peptide sequence as set out in any of SEQ ID NOs.: 25,435 - 37,811, or a complementary sequence or a fragment thereof wherein said probe hybridizes at high stringency to a nucleic acid expressed in the human brain.

16. A single exon nucleic acid probe as claimed in any one of claims 13 to 15 wherein said single exon nucleic acid probe comprises between 15 and 25 contiguous nucleotides of said SEQ ID NO.

17. A single exon nucleic acid probe as claimed in any one of claims 13 to 15, wherein said probe is between 3 - 25 kb in length.

18. A single exon nucleic acid probe as claimed in any one of claims 13 - 17, wherein said probe is DNA, RNA or PNA.

25

19. A single exon nucleic acid probe as claimed in any one of claims 13 - 18, wherein said probe is detectably labeled.

30 20. A single exon nucleic acid probe as claimed in any one of claims 13 - 19, wherein said probe lacks prokaryotic and bacteriophage vector sequence.

35 21. A single exon nucleic acid probe as claimed in any one of claims 13 - 20, wherein said probe lacks homopolymeric

stretches of A or T.

22. A method of measuring gene expression in a sample derived from human brain, comprising:

5        contacting the microarray of claim 12, with a first collection of detectably labeled nucleic acids, said first collection of nucleic acids derived from mRNA of human brain; and then  
10      measuring the label detectably bound to each probe of said microarray.

23. A method of identifying exons in a eukaryotic genome, comprising:

15      algorithmically predicting at least one exon from genomic sequence of said eukaryote; and then detecting specific hybridization of detectably labeled nucleic acids to a single exon probe, wherein said detectably labeled nucleic acids are derived from mRNA from the brain of said eukaryote, said probe is a  
20      single exon probe having a fragment identical in sequence to, or complementary in sequence to, said predicted exon, said probe is included within a microarray according to claim 12, and said fragment is selectively hybridizable at high stringency.

25

24. A method of assigning exons to a single gene, comprising:

30      identifying a plurality of exons from genomic sequence according to the method of claim 23; and then  
measuring the expression of each of said exons in a plurality of tissues and/or cell types using hybridization to single exon microarrays having a probe with said exon,  
35      wherein a common pattern of expression of said exons in

said plurality of tissues and/or cell types indicates that the exons should be assigned to a single gene.

25. A nucleic acid sequence as set out in any of SEQ ID  
5 NOS: 1 - 25,434 which encodes a peptide.
26. A peptide encoded by a sequence as set out in any of SEQ ID Nos: 1 - 25,434.
- 10 27. A peptide comprising a sequence as set out in any of SEQ ID NOS: 25,435 - 37,811.

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 Table 4  
 Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar BLAST E Value	(Top) Hit No.	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
437	13223	25868	6.47					
889	136338	26308	15.92					
1022	13782		2.15					
1279	14029	26898	10.88					
1488	14235	26921	1.22					
1488	14236	26922	1.22					
1609	14355	27044	3.19					
1633	14378	27058	6.1					
1718	14461	27160	3.31					
1743	14485	27184	1.44					
1750	14492	27192	6.78					
1884	14621	27331	1.44					
1871	14707	27425	2.14					
2162	14892	27627	2.7					
2277	15003	27743	2.91					
2578	15292	28028	1					
2578	15292	28028	1					
3181	15944	28595	2.83					
3442	16198	28648	1.42					
3605	16261	28915	12.04					
3549	16304		1					
3649	16402	29042	1.67					
3928	16578		1.03					
4173	16913	29543	1.52					
4230	16971	29595	6.4					
4248	16989	29613	0.97					
4248	16989	29614	0.97					
4303	17042		1.07					
4361	17099	29734	0.78					
4784	17516	30138	0.99					
4983	17708	30310	5.38					
4985	17718	30323	1.3					
5176	17985	30500	1.57					
5176	17985	30501	1.57					

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Table 4

## Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
5336	18139			4.3			
5510	18308			0.14			
5593	18139			3.97			
5848	184443	319398		0.6			
5854	184448	319392		3.28			
5932	25082	319573		1.62			
5958	18740	319699		1.76			
6322	19092			1.27			
6454	19222	32220		1.1			
6454	19222	32221		1.1			
7026	19717	32774		1			
7025	19717	32775		1			
7311	19694	33071		1.78			
7311	19694	33072		1.78			
7712	20376			0.61			
7880	20655	33780		1.4			
8284	21077	34214		1.49			
8758	21451	34598		0.59			
8759	21451	34599		0.59			
9434	22112	35287		2.67			
9668	22318	35515		0.77			
9782	22433	35638		1.24			
8922	22570	35767		0.84			
10328	22975	36194		0.62			
10328	22975	36195		0.62			
10682	23277			2.53			
10749	25151	36879		1.34			
10952	23629			2.22			
11080	23701	36968		1.84			
11132	24023	37328		2.02			
11485	24088			2.47			
12213	24735			1.52			
12609	24916	31006		2.38			
5861	18743	31703		17.79	9.8E+00	AJ239028.1	NT

Homo sapiens LSS gene, partial, exons 15, 16, 17 and 18

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**Table 4**  
**Single Exon Probes Expressed in Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
7905 20600	33730	1.74	9.8E+00	U32718.1	NT		Haemophilus influenzae Rd section 31 of the complete genome
9843 22285	35489	0.44	9.8E+00	Y18930.1	NT		Sulfobdota sofiataracticus 281 kb genomic DNA fragment, strain F2
9843 22285	35490	0.44	9.8E+00	Y18930.1	NT		Sulfobdota sofiataracticus 281 kb genomic DNA fragment, strain F2
6901 19839	32684	0.73	9.6E+00	AF065830.1	NT		Gallus gallus ornithine transcarbamylase (OTC) gene, exon 1
6901 19839	32685	0.73	9.6E+00	AF065830.1	NT		Gallus gallus ornithine transcarbamylase (OTC) gene, exon 1
10321 22968	36187	1.17	9.6E+00	AF242432.1	NT		Mus musculus Nap3 gene, exon 1; neuronal apoptosis inhibitory protein 1 (Nrip1) and general transcription factor IIH polypeptide 2 (Grif2h2) genes, complete cds
10321 22968	36188	1.17	9.6E+00	AF242432.1	NT		Mus musculus Nap3 gene, exon 1; neuronal apoptosis inhibitory protein 1 (Nrip1) and general transcription factor IIH polypeptide 2 (Grif2h2) genes, complete cds
2871 15381	28118	1	9.4E+00	L111433.1	NT		Dengue virus type 3 membrane protein (prM/M)/envelope glycoprotein (E) polyprotein mRNA, partial cds
2871 15381	28120	1	9.4E+00	L111433.1	NT		Dengue virus type 3 membrane protein (prM/M)/envelope glycoprotein (E) polyprotein mRNA, partial cds
2824 16690	28334	2.87	9.4E+00	AB043785.1	NT		Mus musculus A173 gene for antithrombin, complete cds
7897 20692	33820	0.91	9.3E+00	AF130980.1	NT		Homo sapiens ectodysplasin-A receptor protein (EDAR) gene, exons 2, 3, and 4
8901 21552	34733	3.06	9.3E+00	P11210	SWISSPROT		IMMEDIATE-EARLY PROTEIN 1 (IE1) IMMEDIATE-EARLY PHOSPHOPROTEIN PP89
5214 18022	30845	2.46	9.1E+00	AF095809.1	NT		Leuciscus cephalus orientalis cytochrome b (cyt b) gene, partial cds; mitochondrial gene for mitochondrial product
5214 18022	30846	2.46	9.1E+00	AF095809.1	NT		Leuciscus cephalus orientalis cytochrome b (cyt b) gene, partial cds; mitochondrial gene for mitochondrial product
58330 216977		0.83	9.0E+00	P08241	SWISSPROT		RHODOPSIN
8945 19727	31685	5.55	8.9E+00	BE071808.1	EST_HUMAN		601651038R1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:3834592 3'
62287 18060	32041	2.28	8.7E+00	AB019788.1	NT		Cynops pyrrhogaster Cp1bx3 premature mRNA, partial cds
62287 19060	32042	2.28	8.7E+00	AB019788.1	NT		Cynops pyrrhogaster Cp1bx3 premature mRNA, partial cds
430 13216	25881	2.3	8.4E+00	5031804	NT		Homo sapiens Insulin receptor substrate 1 (IRS1) mRNA
6355 20428	33545	3.98	8.1E+00	AJ131719.1	NT		Zea mays mRNA for legumain-like protease (see2a)
11122 23791		2	8.0E+00	P41820	SWISSPROT		BREFELDIN A RESISTANCE PROTEIN
8051 20745		0.58	7.6E+00	Z21489.1	NT		African swine fever virus NP1450L gene encoding RNA polymerase largest subunit
7246 19931		1.9	7.5E+00	AL448065.1	NT		Theroplasma acidophilum complete genome, segment 3/5
8259 20853	34090	1.61	7.5E+00	P35441	SWISSPROT		THROMBOSPONDIN 1 PRECURSOR
8259 20853	34091	1.61	7.5E+00	P35441	SWISSPROT		THROMBOSPONDIN 1 PRECURSOR
5711 18504	31426	2.66	7.4E+00	BF700517.1	EST_HUMAN		602128876P1 NIH_MGC_68 Homo sapiens cDNA clone IMAGE:4285508 5'
8651 21343	34487	2.7	7.4E+00	P04928	SWISSPROT		HISTIDINE-RICH GLYCOPROTEIN PRECURSOR
8651 21343	34488	2.7	7.4E+00	P04929	SWISSPROT		HISTIDINE-RICH GLYCOPROTEIN PRECURSOR

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Table 4  
Single Exon Probes Expressed In Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
2977	15743	28390	3.58	7.2E+00	L12051.1	NT	Lycopersicum esculentum Mill. GTPase (SAR2) mRNA, complete cds
2977	15743	28391	3.58	7.2E+00	L12051.1	NT	Lycopersicum esculentum Mill. GTPase (SAR2) mRNA, complete cds
6831	19867	32713	0.71	7.2E+00	BE179090.1	EST_HUMAN	RC0-HT0613-200300-031-007 HT0613 Homo sapiens cDNA
7049	19740	32800	1.28	7.1E+00	P28168	SWISSPROT	ZINC-FINGER PROTEIN 1 (ZINC-FINGER HOMEODOMAIN PROTEIN 1)
7049	19740	32801	1.28	7.1E+00	P28168	SWISSPROT	ZINC-FINGER PROTEIN 1 (ZINC-FINGER HOMEODOMAIN PROTEIN 1)
9498	221151		8.63	7.1E+00	AL161685.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 91
11359	24047	37350	3.23	7.1E+00	PD5850	SWISSPROT	HYPOTHETICAL 17.3 kDa PROTEIN IN MRDA-PHPB INTERGENIC REGION
9882	22532	35729	3.37	7.0E+00	P48610	SWISSPROT	ARGININE KINASE (AK)
11215	23878	37165	1.51	7.0E+00	C22469	SWISSPROT	WD-40 REPEAT PROTEIN MS13
8181	20875	34011	1.92	6.9E+00	P35679	SWISSPROT	60S RIBOSOMAL PROTEIN L4 (L2)
10249	22897	36107	1.38	6.9E+00	P44834	SWISSPROT	DNA MISMATCH REPAIR PROTEIN MUTS
10267	22845	36125	0.47	6.9E+00	P34228	SWISSPROT	SKTS PROTEIN
7908	20503	33623	1.53	6.8E+00	WC3412.1	EST_HUMAN	207611.1 Scores melanocyte 2NISHM Homo sapiens cDNA clone IMAGE:2918805
7808	20503	33624	1.53	6.8E+00	WC3412.1	EST_HUMAN	207611.1 Scores melanocyte 2NISHM Homo sapiens cDNA clone IMAGE:2918805
9031	21721		1.20	6.8E+00	P36307	SWISSPROT	OUTER CAPSID PROTEINS VP5 AND VP8
10109	22757	35569	3.24	6.8E+00	Q03570	SWISSPROT	HYPOTHETICAL 15.0 kDa PROTEIN C38C10.5 IN CHROMOSOME III
5202	18010		0.72	6.8E+00	QB9028	SWISSPROT	CATECHOL-O-METHYLTRANSFERASE, SOLUBLE FORM (S-COMT)
6450	19218	32216	0.61	6.8E+00	BF672121.1	EST_HUMAN	602152573EF1 NIH MGIC_81 Homo sapiens cDNA clone IMAGE:2884275
8974	22622	35827	2.36	6.8E+00	Q9ZE07	SWISSPROT	URIDYLATE KINASE (UK) (URIDYL MONOPHOSPHATE KINASE) (UMP KINASE)
8974	22622	35828	2.36	6.8E+00	Q9ZE07	SWISSPROT	URIDYLATE KINASE (UK) (URIDYL MONOPHOSPHATE KINASE) (UMP KINASE)
11073	23143		1.97	6.8E+00	Q10309	SWISSPROT	PROBABLE CATION TRANSPORTING ATPASE C8C3.05C
9078	21788	34931	7	6.8E+00	P03374	SWISSPROT	ENV POLYPROTEIN [CONTAINS: COAT PROTEIN GP62; COAT PROTEIN GP36]
10203	22851	36067	0.49	6.8E+00	BE886001.1	EST_HUMAN	601678435EF1 NIH MGIC_53 Homo sapiens cDNA clone IMAGE:39605895
9842	22284	35488	1.55	6.2E+00	AY010901.1	NT	Schizophyllum commune unknown mRNA
10460	23106	368337	0.5	6.2E+00	6754621	NT	Mus musculus membrane 2, alpha B1 (Man2b1), mRNA
6936	19671	32717	1.48	6.0E+00	BE780163.1	EST_HUMAN	601468034IF1 NIH MGIC_67 Homo sapiens cDNA clone IMAGE:38713035
9718	22367	35565	0.48	6.0E+00	AP0000008.1	NT	Pyrococcus horikoshii OT3 genomic DNA, 1168011-1485000 nt position (6/7)
10411	23057	36274	0.67	6.0E+00	AE001862.1	NT	Deinococcus radiodurans R1 section 1 of 2 of the complete chromosome 2
10411	23057	36275	0.67	6.0E+00	AE001862.1	NT	Deinococcus radiodurans R1 section 1 of 2 of the complete chromosome 2
6428	19168	32163	7.32	5.9E+00	AF155142.1	NT	Mus musculus mitochondrial kinase 3 (Mill3) and two pore domain K+ channel subunit (Kcnk8) genes, complete cds
3514	16270		0.89	5.8E+00	7061567	NT	Homo sapiens DESC1 protein (DESC1), mRNA
7061	19752	32816	0.85	5.7E+00	AF502048.1	NT	Mus musculus immunoglobulin scavenger receptor IgSR mRNA, complete cds

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Table 4  
Single Exon Probes Expressed In Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
7061	19752	32817	0.95	5.7E+00	AF302048.1	NT	Mus musculus immunoglobulin scavenger receptor IgSR mRNA, complete cds
74683	20142		1.13	5.6E+00	P75080	SWISSPROT	DNA POLYMERASE III, ALPHA CHAIN POLC-TYPE (POLIII)
11466	23223	36458	2.59	5.6E+00	Q55276	SWISSPROT	LYCOPENE BETA CYCLASE
6157	18934	31801	0.69	5.5E+00	P47447	SWISSPROT	HEAT-INDUCIBLE TRANSCRIPTION REPRESSOR HRCA
10678	23369		1.28	5.5E+00	AF175425.1	NT	Mus musculus DNA methyltransferase (Dnmt) gene, exons 30, 31, and 32
11464	23221	38455	3.09	5.5E+00	P11880	SWISSPROT	PNEUMOLYSIN (THIOL-ACTIVATED CYTOLYSIN)
6830	19492	32514	1.14	5.4E+00	X02212.1	NT	Chicken alpha-cardiac actin gene
6830	19492	32516	1.14	5.4E+00	X02212.1	NT	Chicken alpha-cardiac actin gene
7769	20486		1.54	5.4E+00	Q81082	SWISSPROT	VITELLOGENIN PRECURSOR (VTG) [CONTAINS: LIPOVITELLIN LV-1N; LIPOVITELLIN LV-1C; LIPOVITELLIN LV-2]
8898	21380	34534	0.83	5.4E+00	P40379	SWISSPROT	REP1 PROTEIN
8898	21380	34535	0.83	5.4E+00	P40379	SWISSPROT	REP1 PROTEIN
9836	22584	35784	1.83	5.4E+00	Q17084	SWISSPROT	RHODOPSIN
9836	22584	35785	1.83	5.4E+00	Q17084	SWISSPROT	RHODOPSIN
4734	17458	30102	1.32	5.3E+00	L43126.1	NT	Bovine immunodeficiency-like virus surface envelope gene, 5' end of cds
7978	20193		3.23	5.3E+00	P54098	SWISSPROT	DNA POLYMERASE GAMMA (MITOCHONDRIAL DNA POLYMERASE CATALYTIC SUBUNIT)
8882	21573	0.49	5.3E+00	AB03490.1	NT	[Homo sapiens HERPUD1 gene for stress protein Herp, complete cds	
11628	24225	37543	3.2	5.3E+00	Q27805	SWISSPROT	PROTABLE ANTIBACTERIAL PEPTIDE POLYPROTEIN PRECURSOR
8377	18177		0.91	5.2E+00	BE184840.1	EST_HUMAN	QV4-HT0891-270400-118-109 HT0891 Homo sapiens cDNA
10271	22819		0.95	5.2E+00	AF248070.1	NT	Drosophila orientacea R1B retrotransposable element reverse transcriptase gene, partial cds
11150	23817		2	5.2E+00	Q10138	SWISSPROT	HYPOTHETICAL 61.1KD PROTEIN C23E2.03C IN CHROMOSOME 1
8881	21652	34626	0.9	5.1E+00	O16006	SWISSPROT	RHODOPSIN
9725	22376	35577	1.19	5.1E+00	P08182	SWISSPROT	COLICIN N IMMUNITY PROTEIN (MICROCIN N IMMUNITY PROTEIN)
6163	18969	31944	0.72	5.0E+00	BF510443.1	EST_HUMAN	601894910F1 NIH MGCG_19 Homo sapiens cDNA clone IMAGE:4124114_5'
10084	22742		0.69	5.0E+00	BF508651.1	EST_HUMAN	6018940420F1 NIH MGCG_17 Homo sapiens cDNA clone IMAGE:4131500_5'
10330	22877	36187	3.07	5.0E+00	AF162445.2	NT	Canis familiaris skeletal muscle chloride channel ClC-1 (CLCN1) mRNA, complete cds
11260	23922	37214	8.95	5.0E+00	Z833860.1	NT	Mycobacterium tuberculosis H37Rv complete genome; segment 103/182
10132	22780		0.71	4.9E+00	U91328.1	NT	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HLA-H) gene, R-Ret gene, and sodium phosphate transporter (NPT3) gene, complete cds
4039	16784		10.88	4.8E+00	AF185225.1	NT	Eunice straussi histone H3 (HS) gene, partial cds
8054	20748	33878	0.47	4.8E+00	BF367909.1	EST_HUMAN	RC3-GN0042-100800-011-c10 GN0042 Homo sapiens cDNA
8429	21131		5.23	4.8E+00	AW759087.1	EST_HUMAN	PM0-BT0547-310100-002-b04 BT0547 Homo sapiens cDNA
283	13080	25731	1.86	4.7E+00	BF240552.1	EST_HUMAN	601875654F1 NIH MGCG_55 Homo sapiens cDNA clone IMAGE:40897716_5'

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Table 4

Single Exon Probes Expressed in Brain

Probe Seq ID No:	Exon Seq ID No:	ORF Seq ID No:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
284	13090	25731	1.89	4.7E+00	BF240552.1	EST_HUMAN	601875634F1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:4089716 5'
3268	16030	28679	2.38	4.7E+00	AL163280.2	NT	Homo sapiens chromosome 21 segment HS21C380
8085	21783	34948	1.18	4.6E+00	BE046437.1	EST_HUMAN	7688g10_x1 NCI_CGAP_CLL1 Homo sapiens cDNA clone IMAGE:32822088 3' similar to TR:O75140 O75140 KIAA0845 PROTEIN. contains element PTR5 repetitive element;
9095	21783	34949	1.18	4.6E+00	BE046437.1	EST_HUMAN	7688g10_x1 NCI_CGAP_CLL1 Homo sapiens cDNA clone IMAGE:32822088 3' similar to TR:O75140 O75140 KIAA0845 PROTEIN. contains element PTR5 repetitive element;
10237	22835		0.61	4.6E+00	AF240786.1	NT	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) genes, complete cds
11054	23724		2.31	4.6E+00	DS9989.1	NT	Synaptobrevin sp. PC028603 complete genome, 18/27 2287260-23892728
11605	24204	37526	2.69	4.5E+00	AE001044.1	NT	Arthaeoglobus fulgidus section 63 of 172 of the complete genome
11782	24353	37685	1.78	4.5E+00	BF0668841.1	EST_HUMAN	602123238F1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4280216 5'
30325	15801	28447	0.96	4.4E+00	BF530893.1	EST_HUMAN	602072585F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:1215284 5'
30325	15801	28448	0.96	4.4E+00	BF530893.1	EST_HUMAN	602072585F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:1215284 5'
6109	18886		1.86	4.4E+00	X13414.1	NT	Murine Ig gene for MHC class I(a) associated invariant chain
60227	18807		0.68	4.3E+00	AF059878.1	NT	Homo sapiens neutrophil collagenase (CLGNA) gene, promoter region and 5UTR
72338	20019	33097	2.03	4.3E+00	Y13402.1	NT	Plasmid vector pR28R+var1 gene, exon 1
75116	20188	33280	0.65	4.3E+00	AE001222.1	NT	Tropenema pallidum section 38 of 91 of the complete genome
10769	23453	36696	7.84	4.3E+00	AF240788.1	NT	Homo sapiens glutathione S-transferase theta 2 (GSTT2) and glutathione S-transferase theta 1 (GSTT1) genes, complete cds
5430	18229		3.44	4.2E+00	P16444	SWISSPROT	MICROSOMAL DIPEPTIDASE PRECURSOR (MDP) (DEHYDROPEPTIDASE-1) (RENAL DIPEPTIDASE) (RDP)
5907	18305	31206	0.87	4.2E+00	P518228	SWISSPROT	LAF-4 PROTEIN (LYMPHOID NUCLEAR PROTEIN)
6874	18891	32627	2.62	4.2E+00	P19883	SWISSPROT	EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN)
6874	18591	32628	2.62	4.2E+00	P19883	SWISSPROT	EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTEIN)
88559	21550	34697	4.68	4.2E+00	AB050913.1	EST_HUMAN	wf87903.x1 Scores: NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2286862 3'
98118	22469	35672	1.06	4.2E+00	P31368	SWISSPROT	NUBBIN PROTEIN (TWAIN PROTEIN) (POU DOMAIN PROTEIN 1) (PDM4-1) (POU-1B) (DOCT1)
10449	22697		0.46	4.2E+00	P40886	SWISSPROT	HEXOSE TRANSPORTER HXT8
5846	25078	31569	0.56	4.1E+00	O09185	SWISSPROT	CELLULAR TUMOR ANTIGEN P63
5846	25079	31570	0.56	4.1E+00	O09185	SWISSPROT	CELLULAR TUMOR ANTIGEN P63
7012	19704	32780	0.84	4.1E+00	BE253688.1	EST_HUMAN	601110727F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3351534 5'
7111	19789	32863	0.86	4.1E+00	BF247889.1	EST_HUMAN	601859035F1 NIH_MGC_38 Homo sapiens cDNA clone IMAGE:4069758 5'
7559	20220	33332	8.73	4.1E+00	O23810	SWISSPROT	YY1 PROTEIN PRECURSOR
7831	20345		0.62	4.1E+00	AB041523.1	NT	Patinoprotein yessoensis mRNA for cathepsin A, complete cds
7883	20347	33459	4.32	4.1E+00	P28984	SWISSPROT	GENE 68 PROTEIN

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Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Meet Similar BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
7683	20347	33480		4.32	4.1E+00 P28984	SWISSPROT	GENE 68 PROTEIN
7817	20512	33638		2.63	4.1E+00 U57503.1	NT	Pan troglodytes novaezebrae LTR element in the RNU2 locus
8440	22118	35295		0.57	4.1E+00 P11263	SWISSPROT	50S RIBOSOMAL PROTEIN L4
8571	22224	35409		2.49	4.1E+00 BF692425.1	EST_HUMAN	602247938FT NIH_MGC_82 Homo sapiens cDNA clone IMAGE:4333209.6 CYCLIN-DEPENDENT KINASE INHIBITOR 1B (CYCLIN-DEPENDENT KINASE INHIBITOR P27) (P27KIP1)
10205	22853			0.48	4.1E+00 P48414	SWISSPROT	HYPOTHETICAL PROTEIN HVLF1
10800	23483			3.06	4.1E+00 P09718	SWISSPROT	601507510F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3909051.5 GLC7-INTERACTING PROTEIN 1
10892	23572			11.69	4.1E+00 BE838880.1	EST_HUMAN	601507510F1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3909051.5 GLC7-INTERACTING PROTEIN 1
36533	16289			0.95	4.0E+00 P38298	SWISSPROT	SUCRASE-ISOMALTASE, INTESTINAL [CONTAINS: SUCRASE : ISOMALTASE]
5372	18600	32524		0.77	4.0E+00 Q62683	SWISSPROT	SUCRASE-ISOMALTASE, INTESTINAL [CONTAINS: SUCRASE : ISOMALTASE]
5372	18600	32525		0.77	4.0E+00 Q62683	SWISSPROT	SUCRASE-ISOMALTASE, INTESTINAL [CONTAINS: SUCRASE : ISOMALTASE]
6838	18600	32524		0.75	4.0E+00 Q62683	SWISSPROT	SUCRASE-ISOMALTASE, INTESTINAL [CONTAINS: SUCRASE : ISOMALTASE]
6838	18600	32525		0.75	4.0E+00 Q62683	SWISSPROT	SUCRASE-ISOMALTASE, INTESTINAL [CONTAINS: SUCRASE : ISOMALTASE]
7089	19778	32843		1.44	4.0E+00 Q33010	SWISSPROT	CELL DIVISION PROTEIN FTSY HOMOLOG
8772	21464	34811		0.45	4.0E+00 Q14157	SWISSPROT	HYPOTHETICAL PROTEIN KIAA0144
9843	2294	36695		0.44	4.0E+00 Q61309	SWISSPROT	NITRIC-OXIDE SYNTHASE (NOS, TYPE I) (NEURONAL NOS) (N-NOS) (MNOS)
10065	22713	35931		0.63	4.0E+00 AE002182.1	NT	Ureaplasma urealyticum section 33 of 59 of the complete genome
11453	23220	36454		1.53	4.0E+00 P14546	SWISSPROT	CYTOCROME C OXIDASE POLYPEPTIDE III
11637	24137	37444		2.27	4.0E+00 P07564	SWISSPROT	GENOME POLYPROTEIN [CONTAINS: CAPSID PROTEIN C (CORE PROTEIN); MATRIX PROTEIN (ENVELOPE GLYCOPROTEIN M); MAJOR ENVELOPE PROTEIN E; NONSTRUCTURAL PROTEINS NS1, NS2A, NS2B, NS4A AND NS4B; RNA-DIRECTED RNA POLYMERASE (NS5)]
11537							GENOME POLYPROTEIN [CONTAINS: CAPSID PROTEIN C (CORE PROTEIN); MATRIX PROTEIN (ENVELOPE GLYCOPROTEIN M); MAJOR ENVELOPE PROTEIN E; NONSTRUCTURAL PROTEINS NS1, NS2A, NS2B, NS4A AND NS4B; HELICASE (NS3); RNA-DIRECTED RNA POLYMERASE (NS5)]
13494	16250	37445		2.27	4.0E+00 P07564	SWISSPROT	N.tabacum chitinase gene 50 for class I chitinase C
4287	17026	26904		4.61	3.9E+00 XG4518.1	NT	Mus musculus seminal vesicle secretory protein 88 (MSVSP88) genes, promoter region
6572	18369	31279		8.24	3.8E+00 AF055468.1	NT	MRO-BN0070-300500-02B-h05 BN0070 Homo sapiens cDNA
5572	18369	31280		2.91	3.8E+00 BE814357.1	EST_HUMAN	MRO-BN0070-300500-02B-h05 BN0070 Homo sapiens cDNA
6591	18364	32367		0.55	3.8E+00 U81328.1	NT	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphate transporter (NPT3) gene, complete cds
6774	19518	32546		4.62	3.9E+00 P35298	SWISSPROT	HYPOTHETICAL TRANSCRIPTIONAL REGULATOR IN AIDS-RPSF INTERGENIC REGION

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**Table 4**  
**Single Exon Probes Expressed in Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
7282	16946	33022	4.3	3.9E+00	M23907.1	NT	Human MHC class II lymphocyte antigen (DPw4-beta-1) gene, exon 2
8216	20910	34048	1.88	3.8E+00	X65865.1	NT	Xlaevis mRNA for M4 muscarinic receptor
11385	23178	36403	3.3	3.9E+00	Y18000.1	NT	Homo sapiens NF2 gene
2636	15347		0.9	3.8E+00	AE001662.1	NT	Helicobacter pylori, strain 199 section 123 of 132 of the complete genome
6297	19070	32054	0.88	3.8E+00	Q57830	SWISSPROT	HYPOTHETICAL PROTEIN MJ0385
6673	19590	32626	0.88	3.8E+00	AI469849.1	EST_HUMAN	Q25107.21 NCI_OGAP_Kid11 Homo sapiens cDNA clone IMAGE:2030437.3'
8331	21024	34161	1.1	3.8E+00	D44725.1	EST_HUMAN	HUMSUPYY135 Human brain cDNA Homo sapiens cDNA clone 148
8684	22345		0.82	3.8E+00	AJ380961.1	NT	Streptococcus oralis partial xpt gene for xanthine phosphoribosyltransferase, strain NCTC7884
4001	16748	28379	12.28	3.7E+00	AL161539.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 39
7088	19757		1.03	3.7E+00	AL445065.1	NT	Thermoplasma acidophilum complete genome; segment 3/5
8609	21301		0.55	3.7E+00	4503950	NT	Homo sapiens glucokinase (hexokinase 4, maturity onset diabetes of the young 2) (GCK), nuclear gene encoding mitochondrial protein, mRNA
9078	21765	34828	0.7	3.7E+00	U43543.1	NT	Mus musculus laminin beta 2 gene, exons 17-38, and complete cds
11408	24057	37363	2.23	3.7E+00	BF0689279.1	EST_HUMAN	802120551F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4277748.5'
11408	24057	37364	2.23	3.7E+00	BF0689279.1	EST_HUMAN	802120551F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4277748.5'
579	13359	25988	5.19	3.8E+00	AV761055.1	EST_HUMAN	AV761055 MDS Homo sapiens cDNA clone MDSBUE10.5'
4746	17477		1.06	3.8E+00	AL161472.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 2
5174	17833	30498	0.74	3.8E+00	BF316816.1	EST_HUMAN	801601881F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:4131016.5'
8450	21142	34280	0.85	3.8E+00	D12367.1	EST_HUMAN	HUM0001TB08 Liver HepG2 cell line, Homo sapiens cDNA clone b008
8450	21142	34281	0.85	3.8E+00	D12367.1	EST_HUMAN	HUM0001TB08 Liver HepG2 cell line, Homo sapiens cDNA clone b008
8543	21235	34978	3.83	3.8E+00	AE00447.1	NT	Pseudomonas aeruginosa PA01, section 8 of 529 of the complete genome
8543	21235	34379	3.83	3.8E+00	AE00447.1	NT	Pseudomonas aeruginosa PA01, section 8 of 529 of the complete genome
10759	23444					NT	Escherichia coli glycerophosphate dehydrogenase (gpd) gene, partial cds, and the translation start site has been verified (glpG), and repressor protein (glpR) genes, complete cds
3241	16003	28652	4.07	3.8E+00	M96786.1	NT	Cryptosporidium felis heat shock protein 70 (HSP70) gene, partial cds
5911	18695		1.1	3.8E+00	AF221538.1	NT	Bonellia burdettorum (strain 25015) outer surface protein (ospC) gene, partial cds
6118	18896	31864	1.17	3.8E+00	L42898.1	EST_HUMAN	J940098.11 Severe infant brain TNB Homo sapiens cDNA clone IMAGE:34840.5'
6383	21076		1.18	3.8E+00	R19745.1	SWISSPROT	THROMBOXANE A SYNTHASE (TXA SYNTHASE) (TXS)
8830	21621	34763	1.02	3.8E+00	AA160988.1	EST_HUMAN	ZP86804.51 Strategene HeLa cell s3 837/216 Homo sapiens cDNA clone IMAGE:627055.3' similar to contains Alu repetitive element; contains element MSR1 repetitive element;
8930	21621	34764	1.02	3.8E+00	AA160988.1	EST_HUMAN	ZP86804.51 Strategene HeLa cell s3 837/216 Homo sapiens cDNA clone IMAGE:627055.3' similar to contains Alu repetitive element; contains element MSR1 repetitive element;
8383	22055	35227	0.88	3.8E+00	AL161553.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 33

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Table 4

## Single Exon Probes Expressed In Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
10417	23063	36283	0.46	3.5E+00	AJ33723.1	NT	Bos taurus mRNA for Ran-binding protein 2, partial
1501	14247	26933	2.94	3.4E+00	AF254577.1	NT	Brassica napus RPB5d mRNA, complete cds
7261	18845	33021	2.64	3.4E+00	P04052	SWISSPROT	DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT
7601	20267	33374	0.69	3.4E+00	P04052	SWISSPROT	DNA-DIRECTED RNA POLYMERASE II LARGEST SUBUNIT
8577	21289		0.7	3.4E+00	U65406.1	NT	Human alternatively spliced potassium channels ROM-K1, ROM-K2, ROM-K3, ROM-K4, ROM-K5, and ROM-K6 (KCNJ1) gene, complete cds
8972	21682	34813	0.67	3.4E+00	AJ228042.1	NT	Human sapiens S68 kb contig between AML1 and CBR1 on chromosome 21q22, segment 23
9010	21700	34850	0.54	3.4E+00	AJ250567.1	NT	Human sapiens partial TMASF2 gene for tetraspanin protein, exon 6
10164	22812	36030	2.97	3.4E+00	AF013167.1	NT	Saccharomyces cerevisiae MSS1 gene, complete cds
11619	24119	37429	1.89	3.4E+00	L77570.1	NT	Human sapiens DiGeorge syndrome critical region, centromeric end
5977	18769	31722	1.57	3.3E+00	Q109689	SWISSPROT	PUTATIVE IRON ALCOHOL DEHYDROGENASE
5977	18759	31723	1.57	3.3E+00	Q109689	SWISSPROT	PUTATIVE IRON ALCOHOL DEHYDROGENASE
7784	20469	33611	0.79	3.3E+00	AF111168.2	NT	Human sapiens serine palmitoyl transferase, subunit II gene, complete cds; and unknown genes
10361	23008	36223	0.9	3.3E+00	AP001611.1	NT	Bacillus halodurans genomic DNA, section 5/14
10361	23008	36224	0.9	3.3E+00	AP001611.1	NT	Bacillus halodurans genomic DNA, section 5/14
4883	13273	25908	1.84	3.2E+00	X88422.1	NT	Drosophila sp-50 POU gene
4004	13273	25908	0.9	3.2E+00	X88422.1	NT	Drosophila sp-50 POU gene
4879	17413	30048	1.08	3.2E+00			Human sapiens carboanhydrase antigen-related cell adhesion molecule 1 (biliary glycoprotein) (CEACAM1), mRNA
5481	18280	31178	1.06	3.2E+00	P64924	4502404 NT	SQUALENE-HOPENE CYCLASE, mRNA
5481	18280	31177	1.06	3.2E+00	P64924	SWISSPROT	SQUALENE-HOPENE CYCLASE
5515	18313	31214	2.7	3.2E+00	P12783	SWISSPROT	PHOSPHOGLYCERATE KINASE, CYTOSOLIC
5515	18313	31215	2.7	3.2E+00	P12783	SWISSPROT	PHOSPHOGLYCERATE KINASE, CYTOSOLIC
6214	18988	31964	1.78	3.2E+00	P18834	SWISSPROT	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4
6214	18988	31965	1.78	3.2E+00	P18834	SWISSPROT	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4
7505	20116	33270	0.7	3.2E+00	P04275	SWISSPROT	VON WILLEBRAND FACTOR PRECURSOR (VWF)
7672	20358	33448	2.65	3.2E+00	Y13855.1	NT	Chlamydomonas reinhardtii chloroplast DNA for rps9, ycf4, ycf3, rps18 genes
7672	20358	33449	2.65	3.2E+00	Y13855.1	NT	Chlamydomonas reinhardtii chloroplast DNA for rps9, ycf4, ycf3, rps18 genes
89228	21619		4.61	3.2E+00	P13081	SWISSPROT	PERIPLASMIC [NIFE] HYDROGENASE SMALL SUBUNIT (NIFE HYDROGENASE SMALL CHAIN)
9430	22108	35283	0.87	3.2E+00	M36383.1	NT	S.cerevisea threonine deaminase (ILV1) gene, complete cds
10041	22889	36907	2.03	3.2E+00	AB016081.2	NT	Oryza sativa latifolia OGCA gene for guanylyl cyclase C, complete cds
11946	245010		2.44	3.2E+00	L33836.1	NT	Sus scrofa choline acetyltransferase gene, promoter region
5785	18576	31505	2.46	3.1E+00	Q10135	SWISSPROT	HYPOTHETICAL 142.5 KD PROTEIN C23E2.02 IN CHROMOSOME 1
7287	18970	33047	0.93	3.1E+00	P62178	SWISSPROT	TRIOSE PHOSPHATE TRANSLOCATOR, NON-GREEN PLASTID PRECURSOR (CPT)

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Table 4  
Single Exon Probes Expressed In Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
7627	20293			0.94	3.1E+00 AF303225.1	NT	<i>Bacillus ecelophillus</i> peptidase gene (pefE) gene, complete cds
7985	20681	33807	0.48	3.1E+00 P40885	SWISSPROT	PROBABLE UBIQUITIN-PROTEIN LIGASE HUL4	
8500	21192	34333	4.38	3.1E+00 P48894	SWISSPROT	TYPE IODOTHYRONINE DEIODINASE (TYPE-I 5'DEIODINASE) (DIO1) (TYPE 1 DI) (SDI)	
8500	21192	34334	4.38	3.1E+00 P48894	SWISSPROT	TYPE IODOTHYRONINE DEIODINASE (TYPE-I 5'DEIODINASE) (DIO1) (TYPE 1 DI) (SDI)	
9168	21689		3.85	3.1E+00 Q14957	SWISSPROT	GLUTAMATE [NMDA] RECEPTOR SUBUNIT EPSILON 3 PRECURSOR (N-METHYL-D-ASPARTATE RECEPTOR SUBTYPE 2C) (NR2C) (NMNDAR2C)	
9796	22447	35632	0.58	3.1E+00 7524759	NT	<i>Chloris vulgaris</i> chloroplast, complete genome	
9888	22538		0.83	3.1E+00 Q10125	SWISSPROT	HYPOTHETICAL 56.3 kD PROTEIN F52C9.5 IN CHROMOSOME III DEOXYHYDROSYNTHASE (DHS)	
10234	22882	36035	5.52	3.1E+00 P49385	SWISSPROT	GENOME POLYPROTEIN [CONTAINS: CAPSID PROTEIN C (CORE PROTEIN); MATRIX PROTEIN (ENVELOPE PROTEIN M); MAJOR ENVELOPE PROTEIN E; NONSTRUCTURAL PROTEINS NS1, NS2A, NS2B, NS4A AND NS4B; HELICASE (NS3); RNA-DIRECTED RNA POLYMERASE (NS5)]	
11440	23207		2.68	3.1E+00 P333515	SWISSPROT	retinoic acid nuclear receptor isoform beta 2 [mice, embryonal carcinoma cell line, PCC74A21, mRNA, 2071 nt]	
11443	24006		3.28	3.1E+00 S56680.1	NT	<i>Hamo saurus</i> hypothetical protein PRO0889 (PRO0889), mRNA	
2842	15610	28259	1.09	3.0E+00 8923984	NT	<i>S. aureus</i> genes encoding Sau3A1 DNA methyltransferase and Saug861 restriction endonuclease	
5254	18080	31089	1.32	3.0E+00 X53098.1	NT	Conryobacterium glutamicum trtC gene for threonine synthase (EC 4.2.89.2)	
6461	19228	32228	0.83	3.0E+00 X56037.1	NT	Conryobacterium glutamicum trtC gene for threonine synthase (EC 4.2.89.2)	
8461	19228	32229	0.83	3.0E+00 X56037.1	NT	CYR61 PROTEIN PRECURSOR (3CH61)	
7055	19748		9.09	3.0E+00 P18406	SWISSPROT	ENDOTHELIAL CELL MULTIMERIN PRECURSOR	
7096	19755		0.8	3.0E+00 Q13201	SWISSPROT	B.napus DNA for myosinase	
8805	21497		1.2	3.0E+00 X87838.1	NT	S-ADENOSYLMETHIONINE SYNTHETASE (METHIONINE ADENOSYL TRANSFERASE) (ADOMET SYNTHETASE)	
10182	22840	36055	0.62	3.0E+00 Q58806	SWISSPROT	CD10 PROTEIN HOMOLOG	
10544	23240	36474	1.57	3.0E+00 Q16181	SWISSPROT	RETINAL GUANYLYL CYCLASE 2 PRECURSOR (GUANYLATE CYCLASE 2F, RETINAL) (RETGC2) (ROD OUTER SEGMENT MEMBRANE GUANYLATE CYCLASE 2) (ROS-GC2) (GUANYLATE CYCLASE F) (GC-F)	
10831	23811	36880	0.44	3.0E+00 P51842	SWISSPROT	RETINAL GUANYLYL CYCLASE 2 PRECURSOR (GUANYLATE CYCLASE 2F, RETINAL) (RETGC2) (ROD OUTER SEGMENT MEMBRANE GUANYLATE CYCLASE 2) (ROS-GC2) (GUANYLATE CYCLASE F) (GC-F)	
10831	23811				SWISSPROT	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 4	
11578	24177				SWISSPROT	Chlamydomonas reinhardtii AR39, section 53 of 84 of the complete genome	
2004	14740	27464	2.28	2.9E+00 AE002225.2	NT	F-pringle gidaPA gene for P-protein of the glycine cleavage system	
6809	19470	32483	1.74	2.9E+00 Z36879.1	NT	BRAIN-SPECIFIC ANGIOGENESIS INHIBITOR 1 PRECURSOR	
7110	19798	32861	6.21	2.9E+00 O14814	SWISSPROT		

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**Single Exon Probes Expressed in Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
7110	19788	32882	5.21	2.8E+00 O1454	SWISSPROT	BRAIN-SPECIFIC ANGIOGENESIS INHIBITOR 1 PRECURSOR	
7356	20837	33115	6.84	2.8E+00 P46589	SWISSPROT	ADHERENCE FACTOR (ADHESSION AND AGGREGATION MEDIATING SURFACE ANTIGEN)	
7787	20463	33587	0.67	2.8E+00 P05844	SWISSPROT	STRUCTURAL POLYPROTEIN [CONTAINS: MAJOR STRUCTURAL PROTEIN VP2; NONSTRUCTURAL PROTEIN VP4; MINOR STRUCTURAL PROTEIN VP3]	
7787	20463	33588	0.67	2.9E+00 P05844	SWISSPROT	STRUCTURAL POLYPROTEIN [CONTAINS: MAJOR STRUCTURAL PROTEIN VP2; NONSTRUCTURAL PROTEIN VP4; MINOR STRUCTURAL PROTEIN VP3]	
7886	20691	33819	1.03	2.9E+00 BF344171.1	EST_HUMAN	6020917413F1 NCI_CGAP_Bm64 Homo sapiens cDNA clone IMAGE:4163059 6'	
1440	14167	26872	4.4	2.8E+00 AF186398.1	NT	Buxus harlandii mature K (male) gene, partial cds; chloroplast product	
1629	14276		2.74	2.8E+00 AL161552.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 52	
7207	19892	32888	6.72	2.8E+00 8393724	NT	Mus musculus endomucin (LOC53423), mRNA	
9513	22168		0.54	2.8E+00 BE965182.1	EST_HUMAN	601342758F1 NIH_M6C_53 Homo sapiens cDNA clone IMAGE:3684807 6'	
10588	19892	32888	1.32	2.8E+00 8393724	NT	Mus musculus endomucin (LOC53423), mRNA	
224	13039	25672	13.61	2.7E+00 8678306	NT	Mus musculus per-hecamer repeat gene 3 (Phex3), mRNA	
224	13039	25673	13.51	2.7E+00 8678306	NT	Mus musculus per-hecamer repeat gene 3 (Phex3), mRNA	
5484	18233	31154	1.17	2.7E+00 L14005.1	NT	Homo sapiens apoa1 polymorphism Kringle IV gene, exons 1 and 2	
8045	20739		0.6	2.7E+00 U16947.1	NT	Ipomoea purpurea chalcone synthase (CHS3) gene including complete 5'UTR and complete cds	
8887	21558		1.83	2.7E+00 AL116859.1	NT	Bothrops chilensis strain T4 cDNA library under conditions of nitrogen deprivation	
9332	20403	33519	0.73	2.7E+00 AW098191.1	EST_HUMAN	x888612-21 NCI_CGAP_Bm35 Homo sapiens cDNA clone IMAGE:25813744 3' similar to gb:NM7733	
10397	23043		1.75	2.7E+00 BE063527.1	EST_HUMAN	THYMOSIN BETA-4 (HUMAN); CM0-BT0281-0311989-087-104 BT0281 Homo sapiens cDNA	
4626	17391	28994	5.15	2.8E+00 AF068148.1	NT	Mus musculus sphingomyelin kinase (SPHK1b) mRNA, complete cds	
6480	18289	31149	1.68	2.8E+00 87556601	NT	Mus musculus SRY-box containing gene 13 (Sox13), mRNA	
6480	18289	31150	1.68	2.8E+00 87556601	NT	Mus musculus SRY-box containing gene 13 (Sox13), mRNA	
5736	18523		0.59	2.8E+00 Y170822.1	NT	Mycobacterium fortuitum furA II gene	
7454	25424		0.82	2.8E+00 AJ224639.1	NT	Homo sapiens Surf-5 and Surf-6 genes	
7600	20268		6.04	2.8E+00 AF235902.1	NT	Mus musculus SH2-containing inositol 5-phosphatase (Ship) gene, exons 16 through 27, and complete cds	
7858	20853	33773	1.13	2.8E+00 AJ132180.1	NT	feba been necrotic yellow virus C2-Eg gene, isolate Egyptian EV1-83	
7958	20653	33777	1.13	2.8E+00 AJ132180.1	NT	feba been necrotic yellow virus C2-Eg gene, isolate Egyptian EV1-83	
8557	22210	35393	2.83	2.8E+00 AL161540.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 40	
10253	22901		1.67	2.8E+00 9055163	NT	Mus musculus cleavage and polyadenylation specificity factor 3 (Cpsf3), mRNA	
10953	23630	36873	1.32	2.8E+00 AF143675.1	NT	Hantavirus Z10 segment M G1/G2 glycoprotein (Z10) gene, complete cds	
12560	25304		3.17	2.8E+00 11419220	NT	Homo sapiens ATP-binding cassette, sub-family B (MDR/TAP) member 4 (ABCB4), mRNA	
1449	14195	26873	3.73	2.8E+00 AJ271844.1	NT	Aspergillus nidulans recQ gene for DNA helicase, exons 1-4	

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Table 4

## Single Exon Probes Expressed In Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
1448	14195	26879	3.73	2.5E+00	AJ271844.1	NT	Aspergillus nidulans recQ gene for DNA helicase, exons 1-4
5723	18515	31434	2.22	2.5E+00	P13485	SWISSPROT	TEICOIC ACID BIOSYNTHESIS PROTEIN F
5723	18515	31435	2.22	2.5E+00	P13485	SWISSPROT	TEICOIC ACID BIOSYNTHESIS PROTEIN F
6367	18515	31434	1.63	2.5E+00	P13485	SWISSPROT	TEICOIC ACID BIOSYNTHESIS PROTEIN F
6367	18515	31435	1.63	2.5E+00	P13485	SWISSPROT	TEICOIC ACID BIOSYNTHESIS PROTEIN F
6630	18392	32408	0.84	2.5E+00	D30062.1	NT	Vibrio cholerae cytA gene and cytB gene for cholera toxins, complete cds
7659	20323	33431	0.88	2.5E+00	AW040158.1	EST_HUMAN	QV4-F70005-110500-205-007 F10005 Homo sapiens cDNA
7700	20363	33477	0.58	2.5E+00	4802902	NT	Homo sapiens cathein, heavy poly peptide-like 1,(CLTC1) mRNA
9001	21691	34841	1.53	2.5E+00	D50307.1	NT	Rice DNA for adikotase C-1, complete cds
9752	22403	35608	0.67	2.5E+00	BE287788.1	EST_HUMAN	6011757789F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:3531080 5'
11523	24128		1.34	2.5E+00	P40170	SWISSPROT	DNAJ PROTEIN
11943	24498		3.08	2.5E+00	AF289685.1	NT	Mus musculus EIF4H gene, partial cds; LIMK1 gene, complete cds; and ELN gene, partial cds
3012	167778	28428	1.13	2.4E+00	M24282.1	NT	Chicken alpha-3 collagen type VI mRNA, 3' end
4849	17579	30223	6.09	2.4E+00	4503362	NT	Homo sapiens double C2-like domains, alpha (DOC2A) mRNA
5920	18705	31857	4.18	2.4E+00	P02843	SWISSPROT	VITELLOGENIN 1 PRECURSOR (YOLK PROTEIN 1)
7280	18964	33040	0.78	2.4E+00	Bf9877502.1	EST_HUMAN	60221203858F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4278012 5'
7280	18964	33041	0.78	2.4E+00	Bf9877502.1	EST_HUMAN	60221203858F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4278012 5'
8039	20734	33835	2.4	2.4E+00	P26842	SWISSPROT	CD27L RECEPTOR PRECURSOR (T-CELL ACTIVATION ANTIGEN CD27) (T14)
8039	20734	33836	2.4	2.4E+00	P26842	SWISSPROT	CD27L RECEPTOR PRECURSOR (T-CELL ACTIVATION ANTIGEN CD27) (T14)
8110	20804		2.63	2.4E+00	AE001488.1	NT	Helicobacter pylori, strain J96 section 47 of 132 of the complete genome
8549	21241		1.61	2.4E+00	AW875126.1	EST_HUMAN	RC2-PT004-031289-01-005 PT004 Homo sapiens cDNA
8727	21419	34583	7.36	2.4E+00	P24091	SWISSPROT	ENDOCHITINASE B PRECURSOR (CHNB)
8938	22586	35798	2.68	2.4E+00	P13873	SWISSPROT	SKIN GRANULE PROTEIN PRECURSOR
8938	22586	35799	2.59	2.4E+00	P13873	SWISSPROT	SKIN GRANULE PROTEIN PRECURSOR
10007	22655	35883	1.86	2.4E+00	X92611.1	NT	H_sapiens CTGF gene and promoter region
10143	22791		6.65	2.4E+00	P080589	SWISSPROT	XYLOSE KINASE (XYLUOKINASE)
10220	22863	36079	1.62	2.4E+00	BE28702.1	EST_HUMAN	h63108.x1 NCI CGAP Kid11 Homo sapiens cDNA clone IMAGE:3133187 3'
10220	22863	36080	1.62	2.4E+00	BE28702.1	EST_HUMAN	h63108.x1 NCI CGAP Kid11 Homo sapiens cDNA clone IMAGE:3133187 3'
10490	23138	36364	0.87	2.4E+00	Q51481	SWISSPROT	DENITRIFICATION REGULATORY PROTEIN NIRQ
11331	24022	37327	2.16	2.4E+00	AF158632.2	NT	Fragaria x ananassa cytosolic esterolase peroxidase (ApxSC) gene, ApxSC-c allele, complete cds
1231	13980	26880	13.8	2.3E+00	Z48724.1	NT	G.domesticus artificial single chain antibody gene (L3)
4102	16845		1.35	2.3E+00	AJ401081.1	NT	Bos taurus partial cytB gene for cytochrome b

**Table 4**  
**Single Exon Probes Expressed In Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
5744	18338		0.95	2.3E+00	N88245.1	EST_HUMAN	J7340F Human fetal heart Lambda ZAP Express Homo sapiens cDNA clone J7340 6' similar to PROLYL CARBOXYPEPTIDASE
7354	20935	33113	2.47	2.3E+00	6978554	NT	Rattus norvegicus ATPase, Ca++ transporting, ubiquitin (Atq2a3), mRNA
7495	25425		3.07	2.3E+00	P07189	SWISSPROT	MAJOR CENTROMERE AUTOANTIGEN B (CENTROMERE PROTEIN B) (CENP-B)
7679	29343	33455	1.01	2.3E+00	X50265.1	NT	Mus musculus gene homologues coding for DnaK and DnaJ
9003	21698	34848	0.64	2.3E+00	6835317	NT	Polypeptide chainfolding, mitochondrial, complete genome
9088	21757	34919	1.8	2.3E+00	Q11127	SWISSPROT	ALPHA-(1,3)-FUCOSYLTRANSFERASE (GALACTOSIDE 3-L-FUCOSYLTRANSFERASE)
10704	23395	36832	3.83	2.3E+00	Q07078	SWISSPROT	FUCOSYLTRANSFERASE 4 (FUCT-IV)
11782	24373	37703	3.03	2.3E+00	Bf541987.1	EST_HUMAN	ANXIN VII (SYNEVIN)
11782	24373	37704	3.03	2.3E+00	Bf541987.1	EST_HUMAN	6020691121F1 NIH_3T3 cDNA clone IMAGE:40681173 5'
12157	24642	31089	6.84	2.3E+00	BEB85237.1	EST_HUMAN	6020691121F1 NIH_3T3 cDNA clone IMAGE:40681173 5'
3893	16748	28378	0.95	2.2E+00	AF023528.1	NT	601433673F1 NIH_3T3 cDNA clone IMAGE:3918643 5'
4278	17017	29844	5.01	2.2E+00	D67071.1	NT	Magnipore thiosease Class IV chitin synthetase (ch54) gene, complete cds
4278	17017	29845	5.01	2.2E+00	D67071.1	NT	Rat gene for regucalcin, exon1 (non-coding exon)
							Rat gene for regucalcin, exon1 (non-coding exon)
5258	18084	30692	12.73	2.2E+00	Q88307	SWISSPROT	SORTILIN-RELATED RECEPTOR PRECURSOR (SORTING PROTEIN-RELATED RECEPTOR CONTAINING LDLR CLASS A REPEATS) (MSORLA) (SORLA-1) (LOW-DENSITY LIPOPROTEIN RECEPTOR RELATIVE WITH 11 LIGAND-BINDING REPEATS) (LDLR) (LDLR RELATIVE WITH 11 LIGAND-BINDING REPEATS) (LR11) (>)
							SORTILIN-RELATED RECEPTOR PRECURSOR (SORTING PROTEIN-RELATED RECEPTOR CONTAINING LDLR CLASS A REPEATS) (MSORLA) (SORLA-1) (LOW-DENSITY LIPOPROTEIN RECEPTOR RELATIVE WITH 11 LIGAND-BINDING REPEATS) (LDLR) (LDLR RELATIVE WITH 11 LIGAND-BINDING REPEATS) (LR11) (>)
6258	18084	30693	12.73	2.2E+00	Q88307	SWISSPROT	SORTILIN-RELATED RECEPTOR PRECURSOR (SORTING PROTEIN-RELATED RECEPTOR CONTAINING LDLR CLASS A REPEATS) (MSORLA) (SORLA-1) (LOW-DENSITY LIPOPROTEIN RECEPTOR RELATIVE WITH 11 LIGAND-BINDING REPEATS) (LDLR) (LDLR RELATIVE WITH 11 LIGAND-BINDING REPEATS) (LR11) (>)
5763	18554	31478	1.03	2.2E+00	BE327220.1	EST_HUMAN	RC3-C70254-300800-022->08 CT0254 Homo sapiens cDNA
5763	18554	31479	1.03	2.2E+00	BE327220.1	EST_HUMAN	RC3-C70254-300800-022->08 CT0254 Homo sapiens cDNA
5971	18753	31714	0.84	2.2E+00	BE256983.1	EST_HUMAN	600843401171 NIH_3T3 cDNA clone IMAGE:2868777 3'
6261	18035	32010	3.87	2.2E+00	Q03535	SWISSPROT	MINOR VIRION STRUCTURAL PROTEIN MU-2
6502	19287	32269	3.16	2.2E+00	P51458	SWISSPROT	INSULIN-LIKE GROWTH FACTOR II PRECURSOR (IGF-II) (SOMATOMEDIN A)
6881	17838		3.94	2.2E+00	AA594574.1	EST_HUMAN	nt85b02.s1 NCI_CGAP_Co10 Homo sapiens cDNA clone IMAGE:1058379 3'
7217	18802	32975	0.9	2.2E+00	AA137027.1	EST_HUMAN	2a97f04.71 Strategene fetal retina 837202 Homo sapiens cDNA clone IMAGE:586143 5'
7607	20178	33272	19.2	2.2E+00	AA449012.1	EST_HUMAN	2a05g10.71 Seances_total_fetus_Nb21F8_B9 Homo sapiens cDNA clone IMAGE:755834 5'
7689	20267	33365	0.72	2.2E+00	P54918	SWISSPROT	ALANINE RACEMASE
8001	20696	33823	0.68	2.2E+00	BE301560.1	EST_HUMAN	bb1712.x1 NIH_3T3 cDNA clone IMAGE:29683207 3' similar to gb:D45836 Mouse mRNA for nuclear pore-targeting-complex component of (MOUSE)

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Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
8001	20698	33824	0.58	2.2E+00	BE301560.1	EST_HUMAN	b677h12_x1_NIH_MGC_21 Homo sapiens cDNA clone IMAGE:2963207 3' similar to gb:D45836 Mouse mRNA for nuclear pore-targeting-complex component of (MOUSE);
9241	21820		11.02	2.2E+00	BE741978.1	EST_HUMAN	60169473F1 NIH_MGC_9 Homo sapiens cDNA clone IMAGE:3948561 5'
9488	25124		2.28	2.2E+00	Q04706	SWISSPROT	TRANSPOSON TY1 PROTEIN A
9853	22601	35804	1.1	2.2E+00	AI280373.1	EST_HUMAN	qtm68b03_x1_Sorens_placenta_80weeks_2NbHP8e9W Homo sapiens cDNA clone IMAGE:1883985 3' similar to gb:Y00433 GLUTATHIONE PEROXIDASE (HUMAN);
9853	22601	35805	1.1	2.2E+00	AI280373.1	EST_HUMAN	qtm68b03_x1_Sorens_placenta_80weeks_2NbHP8e9W Homo sapiens cDNA clone IMAGE:1883986 3' similar to gb:Y00433 GLUTATHIONE PEROXIDASE (HUMAN);
9896	22644	35858	2.68	2.2E+00	BF246782.1	EST_HUMAN	601855691F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:40753391 5'
- 10353	23000	36217	3.11	2.2E+00	AF183416.1	NT	Homo sapiens ovary granulosa cell 13.0 kDa protein hGR74 homolog mRNA, complete cds
11418	23185	38415	3.47	2.2E+00	P07911	SWISSPROT	UROMODULIN PRECURSOR (TAMM-HORSFALL URINARY GLYCOPROTEIN) (THFP)
11616	24214	37639	5.89	2.2E+00	P10407	SWISSPROT	EARLY1A 28 KD PROTEIN
558	16545	25967	8.3	2.1E+00	AF132612.2	NT	Mus musculus pre-T cell receptor alpha gene, enhancer region and upstream region
3875	16530		1.08	2.1E+00	AW448386.1	EST_HUMAN	U+H-B13-kd-e08-0-1   s1 NCI CGAP_Sub5 Homo sapiens cDNA clone IMAGE:2734850 3'
60441	16821		0.89	2.1E+00	P75357	SWISSPROT	HYPOTHETICAL PROTEIN NG3022_HOMOLOG
6710	19625	32669	3.95	2.1E+00	O70168	SWISSPROT	ALPHA-2 HS-GLYCOPROTEIN PRECURSOR (FETUIN-A)
6946	18428	32443	5.72	2.1E+00	N2875.1	EST_HUMAN	Y08a10_s1_Sorens_melanocyte_2NbHP Homo sapiens cDNA clone IMAGE:270618 3' similar to gb:ME5654 TRANSCRIPTION INITIATION FACTOR TFIID (HUMAN);
83295	21088		1.97	2.1E+00	AU123630.1	EST_HUMAN	AU123630_NT2RM2_Homo sapiens cDNA clone NT2RM20000871 5'
1174	13927	26591	1.44	2.0E+00	AF180527.1	NT	Homo sapiens p22Dokel (DOKDEL) mRNA, complete cds
1174	13927	26592	1.44	2.0E+00	AF180527.1	NT	Homo sapiens p22Dokel (DOKDEL) mRNA, complete cds
1312	14060	26735	0.97	2.0E+00	AF204927.1	NT	Oryctodinus cuniculus $\text{Na}^+$ - $\text{K}^+$ -ATPase beta 1 subunit mRNA, complete cds
1569	14316		2.61	2.0E+00	P25582	SWISSPROT	PUTATIVE RNA METHYLTRANSFERASE SPB1
2145	14875	27609	5.88	2.0E+00	Z78279.1	NT	R.norvegicus mRNA for collagen alpha1 type I
2145	14875	27610	5.88	2.0E+00	Z78279.1	NT	R.norvegicus mRNA for collagen alpha1 type I
4080	16824	28450	2.2	2.0E+00	AW684496.1	EST_HUMAN	h13cg5_x1_NCI CGAP_G11 Homo sapiens cDNA clone IMAGE:2972168 3' similar to gb:J01677 GLYCERALDEHYDE 3-PHOSPHATE DEHYDROGENASE, LIVER (HUMAN);
4080	16824	29451	2.2	2.0E+00	AW684496.1	EST_HUMAN	h13cg5_x1_NCI CGAP_G11 Homo sapiens cDNA clone IMAGE:2972168 3' similar to gb:J01677 GLYCERALDEHYDE 3-PHOSPHATE DEHYDROGENASE, LIVER (HUMAN);
7449	20125		0.92	2.0E+00	P07588	SWISSPROT	STRUCTURAL POLYPROTEIN [CONTAINS: NUCLEOCAPSID PROTEIN C; MEMBRANE GLYCOPROTEINS E1 AND E2]
79223	20618	33745	3.17	2.0E+00	AB008878.1	NT	Escherichia coli 0157 DNA, map position at 46 min., complete cds
79223	20618	33746	3.17	2.0E+00	AB008878.1	NT	Escherichia coli 0157 DNA, map position at 46 min., complete cds
79223	20618	33747	3.17	2.0E+00	AB008878.1	NT	Escherichia coli 0157 DNA, map position at 46 min., complete cds

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**Table 4**  
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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit No.	Top Hit Accession	Top Hit Database Source	Top Hit Descriptor
8820	21512	34656	3.15	2.0E+00	F31500.1	EST_HUMAN	HSPD22703	Hm3 Homo sapiens cDNA clone s+000117B08
12481	26285	30720	7.27	2.0E+00	5834843	NT	Gallus gallus mitochondrial, complete genome	
55111	18309	31209	4.77	1.8E+00	6754389	NT	Mus musculus Inositol 1,4,5-triphosphate receptor 1 (Itp1), mRNA	
55111	18309	31210	4.77	1.8E+00	6754389	NT	Mus musculus Inositol 1,4,5-triphosphate receptor 1 (Itp1), mRNA	
60359	18790	31753	4.32	1.9E+00	BEP69385.1	EST_HUMAN	601678368F1 NIH_MGC_78 Homo sapiens cDNA clone IMAGE:39498815	
65556	19321		0.75	1.9E+00	AW845889.1	EST_HUMAN	MRO-CT0063-071088-002-812 C10063 Homo sapiens cDNA	
68550	18412		2.46	1.9E+00	Q163927	SWISSPROT	CTD-BINDING SR-LIKE PROTEIN RA4	
83258	21051	34190	2.18	1.9E+00	P02487	SWISSPROT	COLLAGEN ALPHABET 2(I) CHAIN PRECURSOR	
83258	21051	34191	2.18	1.9E+00	P02487	SWISSPROT	COLLAGEN ALPHABET 2(I) CHAIN PRECURSOR	
85537	21249		2.94	1.8E+00	BF380206.1	EST_HUMAN	CMB-MTO114-010800-323-H12 M70114 Homo sapiens cDNA	
87922	21484		1.33	1.8E+00	O51781	SWISSPROT	ARGININE DEIMINASE (ADI) (ARGININE DIHYDROLASE) (AD)	
9530	22183	35597	0.59	1.9E+00	AA8689125.1	EST_HUMAN	elb94ed04.57 Stratagene lung (#837210) Homo sapiens cDNA clone IMAGE:8545743 similar to contains Alu repetitive element;contains element L1 L1 repetitive element;	
10462	23108	36539	0.62	1.9E+00	AF248269.1	NT	Homo sapiens gap-gap precursor protein gene, partial cds	
3098	15854	28498	1.13	1.8E+00	P21004	SWISSPROT	PROTEIN B8 PRECURSOR	
							Synechococcus sp. PCG7942 copper transporting P-ATPase (ctaA) and ATP synthase epsilon subunit (atpE) genes, complete cds	
3118	15883	28522	1.57	1.8E+00	U04356.1	NT	Synechococcus sp. PCG7942 copper transporting P-ATPase (ctaA) and ATP synthase epsilon subunit (atpE) genes, complete cds	
5777	18568	28523	1.67	1.8E+00	U04356.1	NT	Synechococcus sp. PCG7942 copper transporting P-ATPase (ctaA) and ATP synthase epsilon subunit (atpE) genes, complete cds	
8013	18794	31757	1.91	1.8E+00	P18502	SWISSPROT	HEDGEHOG RECEPTOR (PATCHED PROTEIN)	
83056	18077		1.32	1.8E+00	BF311998.1	EST_HUMAN	6011887854F1 NIH_MGC_19 Homo sapiens cDNA clone IMAGE:41272645	
8641	19403	32418	1.12	1.8E+00	BF683327.1	EST_HUMAN	602119470F1 NIH_MGC_48 Homo sapiens cDNA clone IMAGE:42982725	
69558	19440	32455	1.64	1.8E+00	BP305652.1	EST_HUMAN	601188348BF1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:41390385	
			1.79	1.8E+00	P21249	SWISSPROT	MAJOR ANTIGEN RETROVIRUS-RELATED POLYPROTEIN [CONTAINS:REVERSE TRANSCRIPTASE ; ENDONUCLEASE]	
8018	20711	33841	0.93	1.8E+00	P11369	SWISSPROT	RETROVIRUS-RELATED POLYPROTEIN [CONTAINS:REVERSE TRANSCRIPTASE ; ENDONUCLEASE]	
8018	20711	33842	0.93	1.8E+00	P11369	SWISSPROT	RETROVIRUS-RELATED POLYPROTEIN [CONTAINS:REVERSE TRANSCRIPTASE ; ENDONUCLEASE]	
83268	21061	34201	0.44	1.8E+00	P48634	SWISSPROT	LARGE PROLINE-RICH PROTEIN BAT2 (HLA-B-ASSOCIATED TRANSCRIPT 2)	
83268	21061	34202	0.44	1.8E+00	P48634	SWISSPROT	LARGE PROLINE-RICH PROTEIN BAT2 (HLA-B-ASSOCIATED TRANSCRIPT 2)	
83268	21061	34203	0.44	1.8E+00	P48634	SWISSPROT	LARGE PROLINE-RICH PROTEIN BAT2 (HLA-B-ASSOCIATED TRANSCRIPT 2)	
87763	21445	34658	1.98	1.8E+00	O43281	SWISSPROT	EMBRYONAL FYN-ASSOCIATED SUBSTRATE (HEFS)	
9073	21762	34924	0.77	1.8E+00	R31042.1	EST_HUMAN	YH72608.1 Soesies placenta Nb2HP Homo sapiens cDNA clone IMAGE:1352785	
9161	21831	34994	0.78	1.8E+00	AW880004.1	EST_HUMAN	QV0-OT0030-070300-148-at3 OT0030 Homo sapiens cDNA	

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Table 4

## Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar BLAST E Value	T <sub>op</sub> Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
9749	22400	35805	0.75	1.8E+00	P27050	SWISSPROT	CHITINASE D PRECURSOR
10183	22831		3.2	1.8E+00	AF111849.1	NT	Homo sapiens PRO0530 mRNA, complete cds
10452	23098		0.63	1.8E+00	P44325	SWISSPROT	CYTIDINE DEAMINASE (CYTIDINE AMINOHYDROLASE) (CDA)
12278	25238		6.29	1.8E+00	AF311284.1	NT	Chlamydomonas reinhardtii alternative oxidase 1 (AOX1) gene, nuclear gene encoding mitochondrial protein
12359	24763		3.9	1.8E+00	8568404	NT	Rattus norvegicus Actin-related protein complex 1b (Ap-1b), mRNA
1086	13844	28502	2.21	1.7E+00	Q60114	SWISSPROT	LEVANSUCRASE (BETA-D-FRUCTOFURANOSYL TRANSFERASE) (SUCROSE 6-FRUCTOSYL TRANSFERASE)
2269	14995	27734	2.29	1.7E+00	AL168280.2	NT	Homo sapiens chromosome 21 segment HS21C080
23172	15034	27833	2.66	1.7E+00	AI141087.1	EST_HUMAN	oz3sh05.x1 Scareas_NIHIMPu_S1 Homo sapiens cDNA clone IMAGE:1678137'3'
4426	17162	29782	0.81	1.7E+00	Q60114	SWISSPROT	LEVANSUCRASE (BETA-D-FRUCTOFURANOSYL TRANSFERASE) (SUCROSE 6-FRUCTOSYL TRANSFERASE)
5525	18523	31223	1.77	1.7E+00	BE063648.1	EST_HUMAN	CMD-BT0282-171/289-127-805 BT0282 Homo sapiens cDNA
5525	18523	31224	1.77	1.7E+00	BE063546.1	EST_HUMAN	CMD-BT0282-171/289-127-805 BT0282 Homo sapiens cDNA
5527	18711	31686	3.28	1.7E+00	Q51TR8	SWISSPROT	COP2 TRANSCRIPTION FACTOR 1 (COUP-TF1) (COUP-TF I)
7118	18806	32871	1.11	1.7E+00	Q03703	SWISSPROT	HYPOTHETICAL 38.0 kD PROTEIN IN CAT2-AMD1 INTERGENIC REGION
7118	19808	32872	1.11	1.7E+00	Q03703	SWISSPROT	HYPOTHETICAL 38.0 kD PROTEIN IN CAT2-AMD1 INTERGENIC REGION
7763	20449	33673	0.91	1.7E+00	AF021385.1	NT	Mus musculus T cell receptor gamma locus, TCR gamma 2 and gamma 4 gene clusters
7832	20827	33755	1.13	1.7E+00	6755715	NT	Mus musculus T-cell acute lymphocytic leukemia 1 (Tal1), mRNA
7861	20658	33781	0.58	1.7E+00	BFS30580.1	EST_HUMAN	602071917F1_NCL CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4214869'5'
8440	21132	34288	0.5	1.7E+00	AE24553.1	NT	Hippoboscidus hippoglossus interferon inducible Mx protein (Mx) mRNA, complete cds
8525	21217		2.3	1.7E+00	BF308000.1	EST_HUMAN	601894255F1_NH_MGC_17 Homo sapiens cDNA clone IMAGE:4140084'5'
8605	21297	34440	0.59	1.7E+00	X690683.1	NT	M.musculus Ank-1 mRNA for erythroid erythrin
8605	21297	34441	0.69	1.7E+00	X690683.1	NT	M.musculus Ank-1 mRNA for erythroid erythrin
9047	25123	34882	2.18	1.7E+00	OE0479	SWISSPROT	HOMEBOX PROTEIN DLX-3
9047	25123	34883	2.18	1.7E+00	OE0479	SWISSPROT	HOMEBOX PROTEIN DLX-3
9506	22160		1.15	1.7E+00	AF161380.1	NT	Homo sapiens HSPC282 mRNA, partial cds
10071	22719		0.48	1.7E+00	AW953681.1	EST_HUMAN	EST366751 MAGE sequences, MAGE Homo sapiens cDNA
11693	24195	37514	2.57	1.7E+00	W22424.1	EST_HUMAN	67B7 Human retina DNA Tsp5084-d cleaved sublibrary Homo sapiens cDNA not directional
12231	24684	31074	1.9	1.7E+00	AI878443.1	EST_HUMAN	MSR1 repetitive element;
12717	24690	30970	1.84	1.7E+00	AI198573.1	EST_HUMAN	qf50b01.x1 Scareas_NHT Homo sapiens cDNA clone IMAGE:1753417'3' similar to contains L1.H1.L1 repetitive element;
2027	14762	27491	18.61	1.6E+00	AF169339.1	NT	Homo sapiens lens epithelium-derived growth factor gene, alternatively spliced, complete cds

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
2036	14771	27500	3.75	1.6E+00	AF077374.1	NT	Homo sapiens small proline-rich protein (SPRR3) gene, exons 1, 2, and 3 and complete cds
2042	14776	27505	1.54	1.6E+00	Y11344.1	NT	Mus musculus ST6GalNAcII gene, exon 2
2282	15007		1.24	1.6E+00	X983373.1	NT	B. <i>napus</i> gene encoding endo-polypeptiduronase zd2ff01_1.1. Scores_fatel_heart_NbH19W Homo sapiens cDNA clone IMAGE:3416895 similar to gb:D288605 N-AcetylLACTOSAMINE SYNTHASE (HUMAN);
2861	15727	28377	1.61	1.6E+00	W68426.1	EST_HUMAN	
4011	16757		5.69	1.6E+00	BF570077.1	EST_HUMAN	60218603571 NIH_MGC_46 Homo sapiens cDNA clone IMAGE:4310591 3'
4319	17058	29632	1.9	1.6E+00	AF155827.1	NT	Homo sapiens proliferation-associated SNF2-like protein (SMARCA6) mRNA, complete cds
4319	17058	29633	1.9	1.6E+00	AF155827.1	NT	Homo sapiens proliferation-associated SNF2-like protein (SMARCA6) mRNA, complete cds
4942	17869	30277	0.84	1.6E+00	AF075394.1	NT	Urothelium chilensis cytochrome c oxidase subunit I (COI) gene, mitochondrial gene encoding mitochondrial protein, partial cds
4942	17869	30278	0.94	1.6E+00	AF075394.1	NT	Urothelium chilensis cytochrome c oxidase subunit I (COI) gene, mitochondrial gene encoding mitochondrial protein, partial cds
5024	17745	30386	2.86	1.6E+00	Y11344.1	NT	Mus musculus ST6GalNAcII gene, exon 2
5024	17745	30387	2.88	1.6E+00	Y11344.1	NT	Mus musculus ST6GalNAcII gene, exon 2
5737	18529		2.16	1.6E+00	LD4808.1	NT	Brachydorid rero M-IC class II DA-hbeta-2F1 gene, 3' end
5823	18612	31643	0.79	1.6E+00	AF005831.1	NT	Homo sapiens transglutaminase type I (Tgase1) gene, promoter region
6378	19147	32146	0.69	1.6E+00	BF380703.1	EST_HUMAN	IL2-JT0073-060600-145-E02 JT0073 Homo sapiens cDNA
6810	18373	32387	1.08	1.6E+00	AW294881.1	EST_HUMAN	UH-B12-ehr-b-04-04-U1.51 NCI_CGAP_Sub4 Homo sapiens cDNA clone IMAGE:2727511 3'
7145	18332	32901	2.73	1.6E+00	BE697267.1	EST_HUMAN	RC0-CT0416-200700-032-010 CT0416 Homo sapiens cDNA
7929	20624		1.19	1.6E+00	Q46378	SWISSPROT	VIRULENCE FACTOR MVIN HOMOLOG
8277	20871	34112	3.28	1.6E+00	AJ2877131.1	NT	Mus musculus SII, MAP_17, CYP_a, SCL & CYP_b genes
8798	21490	34636	0.83	1.6E+00		11437222	NT
8798	21490	34637	0.83	1.6E+00		11437222	NT
8970	21680	34810	0.47	1.6E+00	BE388391.1	EST_HUMAN	601283525F1 NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3605647 5'
8980	25121	33549	1.84	1.6E+00	X52046.1	NT	M. <i>musculus</i> COL3A1 gene for collagen alpha-1
8980	25121	33550	1.84	1.6E+00	X52046.1	NT	M. <i>musculus</i> COL3A1 gene for collagen alpha-1
9487	22140		0.68	1.6E+00	AF043488.1	NT	Thermoanaerobacter ethanolicus D-glycose-binding protein (xyfF) gene, complete cds
9534	22286	35480	1.32	1.6E+00	T41280.1	EST_HUMAN	phb6_18/1TV Outward Alu-primed hnRNA library Homo sapiens cDNA clone pHb6_18/1TV
10047	22685	35911	0.5	1.6E+00	AF121381.1	NT	Drosophila melanogaster signal transducing adapter protein (STAM), serine/threonine kinase Iai (IAI), and zinc finger protein (DNZ1) genes, complete cds
10085	22733	35947	1.15	1.6E+00	AW835844.1	EST_HUMAN	QV4-LT0076-080200-100-d07 LT0018 Homo sapiens cDNA
10085	22733	35948	1.15	1.6E+00	AW835844.1	EST_HUMAN	QV4-LT0076-080200-100-d07 LT0018 Homo sapiens cDNA
10242	22890	36102	0.47	1.6E+00	AF037352.1	NT	Mus musculus T cell receptor gamma locus, TCR gamma 1 and gamma 3 gene clusters
10481	23137	36365	0.45	1.6E+00	AF162084.1	NT	Glucosidase/beta-tubulin 2 (btub2) gene, partial cds

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Table 4

## Single Exon Probes Expressed In Brain

Probe Seq ID No:	Exon Seq ID No:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLASTE Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
10670	23381	36802	1.95	1.8E+00	P54817	SWISSPROT	CAPSID PROTEIN P40 [CONTAINS: ASSEMBLIN (PROTEASE) : CAPSID ASSEMBLY PROTEIN] nc16002_s1 NC_ CGAP_P1 Homo sapiens cDNA clone IMAGE:1008267 similar to contains element MER4 repetitive element;
10728	23416	36857	1.27	1.8E+00	AA216387.1	EST_HUMAN	Homo sapiens transglutaminase type I (Tgase1) gene, promoter region
10747	188112	311643	5.27	1.6E+00	AF056831.1	NT	Rattus norvegicus lun dimerization protein 2 (Ldp-2) mRNA, complete cds
11705	24900	37628	3.46	1.6E+00	AF104313.1	NT	Chlamydomphila pneumoniae AF39, section 32 of 84 of the complete genome
311	12859	25476	5.31	1.5E+00	U53449.1	NT	Mus musculus a disintegrin and metalloprotease domain (ADAM) 15 (metegrin), mRNA
225	13037	25674	2.2	1.5E+00	AE0022201.2	NT	Potato virus A RNA complete genome, isolate U
608	13384		2.03	1.5E+00	8752881	NT	Potato virus A RNA complete genome, isolate U
2410	15131	27867	1.85	1.5E+00	AJ131402.1	NT	Mus musculus T-cell lymphoma invasion and metastasis 1 (Tiam1), mRNA
2519	15235	27976	2	1.5E+00	8878350	NT	Potato virus A RNA complete genome, isolate U
3135	15131	27887	1.85	1.5E+00	AJ131402.1	NT	Defensocidus radiodurans R1 section 82 of 229 of the complete chromosome 1
3368	16127	28785	0.72	1.5E+00	AE01945.1	NT	HT2210_X1 NC_ CGAP_GC3 Homo sapiens cDNA clone IMAGE:2240587 3' similar to TR:O00237 O00237
5842	18437	31350	0.83	1.5E+00	AJ885301.1	EST_HUMAN	HT2210_X1 NC_ CGAP_GC8 Homo sapiens cDNA clone IMAGE:2240587 3' similar to TR:O00237 O00237
5842	18437	31351	0.83	1.5E+00	AJ885301.1	EST_HUMAN	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:1407115 3'
6312	19083	32068	3.02	1.5E+00	R17878.1	EST_HUMAN	HT2210_X1 Severe infant brain NIH Homo sapiens cDNA clone IMAGE:31683 5'
7028	19720		1.37	1.5E+00	BE783356.1	EST_HUMAN	HT2210_X1 Severe NIH MGC_88 Homo sapiens cDNA clone IMAGE:3881555 5'
7060	19751	32814	23.98	1.5E+00	P4779	SWISSPROT	HYPOTHETICAL 11B.4 KD PROTEIN IN BAT2-DAL5 INTERGENIC REGION PRECURSOR
7060	19751	32815	23.98	1.5E+00	P4779	SWISSPROT	HYPOTHETICAL 11B.4 KD PROTEIN IN BAT2-DAL5 INTERGENIC REGION PRECURSOR
7245	18830	331006	0.61	1.5E+00	AA886259.1	EST_HUMAN	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:1407115 3'
7493	20185	33257	0.78	1.5E+00	AJ003254.1	EST_HUMAN	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:1684893 3' similar to HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:3911181 5'
7727	20390		0.84	1.5E+00	AB039887.1	NT	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:3911181 5'
8021	20716	33848	0.89	1.5E+00	BE887448.1	EST_HUMAN	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:4056135 5'
8542	21234	34377	0.84	1.5E+00	K02138.1	NT	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:4056135 5'
8914	21605		0.48	1.5E+00	AB038518.1	NT	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:4056135 5'
9032	21722	34876	0.46	1.5E+00	BF217818.1	EST_HUMAN	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:4056135 5'
8383	22045	35217	0.84	1.5E+00	R81928.1	EST_HUMAN	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:4056135 5'
8635	22188	35374	1.39	1.5E+00	AW378897.1	EST_HUMAN	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:4056135 5'
9750	22411	35518	6.39	1.5E+00	BF376754.1	EST_HUMAN	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:4056135 5'
9952	22600		1.77	1.5E+00	BF337944.1	EST_HUMAN	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:4056135 5'
10096	22744	35658	1.68	1.5E+00	AA017089.1	EST_HUMAN	HT2210_X1 Severe testis NHT Homo sapiens cDNA clone IMAGE:4056135 5'

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Table 4  
Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
10096 22744	35959		1.66	1.5E+00	AA071689.1	EST_HUMAN	2B38608.11 Seires retina N2b4-HR Homo sapiens cDNA clone IMAGE:3813065'
11375 23982	37282		4.46	1.5E+00	AL134197.1	EST_HUMAN	DKFZP547/P243_s1_547 (synonym: hiftr1) Homo sapiens cDNA clone DKFZP547P243_3'
11530 24130		6.55	1.5E+00	X07380.1	NT	Mitro mitochondrial tRNA-Ser gene and tRNA-Pro pseudogene	
11629 24226	37549	2.1	1.5E+00	AI400798.1	EST_HUMAN	Ig34d09.X1 NCI_CGAP_CLL1 Homo sapiens cDNA clone IMAGE:2116433_3'	
11629 24226	37550	2.1	1.5E+00	AI400798.1	EST_HUMAN	Ig34d09.X1 NCI_CGAP_CLL1 Homo sapiens cDNA clone IMAGE:2116433_3'	
12222 25325	30713	1.44	1.5E+00	D83480.1	NT	Human mRNA for KIAA0148 gene, partial cds	
12445 24815		3.38	1.5E+00	AL445035.1	NT	Thermoplasma acidophilum complete genome, segment 3/5	
28 12856	25472	2.76	1.4E+00	7861685	NT	Homo sapiens DKFZP588M0122 protein (DKFZP588M0122), mRNA	
28 12856	25473	2.76	1.4E+00	7861685	NT	Homo sapiens DKFZP588M0122 protein (DKFZP588M0122), mRNA	
2333 15057		6.92	1.4E+00	U67922.1	NT	Ovis aries piton protein gene, complete cds	
2875 16384	28125	2.21	1.4E+00	X74463.1	NT	Human papillomavirus type 7 genomic DNA	
2776 15481	28221	2.61	1.4E+00	AF064564.2	NT	Fugu rubripes neurofibromatosis type 1 (NF1), A-kinase anchor protein (AKAP84), BAW protein (BAW), and WSB1 protein (WSB1) genes, complete cds	
2778 15481	28222	2.61	1.4E+00	AF064564.2	NT	Fugu rubripes neurofibromatosis type 1 (NF1), A-kinase anchor protein (AKAP84), BAW protein (BAW), and WSB1 protein (WSB1) genes, complete cds	
4545 17280		1.81	1.4E+00	BF881547.1	EST_HUMAN	602156837F1 NIH_MGC_83 Homo sapiens cDNA clone IMAGE:4287558_5'	
6298 18053	30754	1.61	1.4E+00	AW054976.1	EST_HUMAN	wk45g07.X1 NCI_CGAP_Pan1 Homo sapiens cDNA clone IMAGE:2510466_3'	
5441 18240		6.57	1.4E+00	AB032983.1	NT	Homo sapiens mRNA for KIAA1157 protein, partial cds	
61863 18963	31838	2.72	1.4E+00	Q13472	SWISSPROT	DNA TOPOISOMERASE II ALPHAI	
62022 25420		4.02	1.4E+00	AB020712.1	NT	Homo sapiens mRNA for KIAA0505 protein, complete cds	
6318 19089	32074	2.67	1.4E+00	Q92777	SWISSPROT	SYNAPSIN II	
6318 19089	32075	2.67	1.4E+00	Q92777	SWISSPROT	SYNAPSIN II	
7186 19872	32646	2.07	1.4E+00	AJ139289.1	NT	Homo sapiens cavinlin-1/-2 locus, Config1, D7S522, genes CAV2 (exons 1, 2a, and 2b), CAV1 (exons 1 and 2)	
7201 19887	32682	1.17	1.4E+00	AW407760.1	EST_HUMAN	Ie230f5.X1 NCI_CGAP_CML1 Homo sapiens cDNA clone IMAGE:28119873_3 similar to contains Ali repetitive element.	
7258 18942	33018	0.75	1.4E+00	P55268	SWISSPROT	LAMININ BETA-2 CHAIN PRECURSOR (S-LAMININ)	
7258 18942	33019	0.75	1.4E+00	P55268	SWISSPROT	LAMININ BETA-2 CHAIN PRECURSOR (S-LAMININ)	
8233 20927		0.68	1.4E+00	P07883	SWISSPROT	GLUCOSAMYLASE PRECURSOR ((GLUCAN 1,4-ALPHA-GLUCOSIDASE)(1,4-ALPHA-D-GLUCAN GLUCOHYDROLASE))	
8683 21385		4.47	1.4E+00	AJ271735.1	NT	Homo sapiens Xq pseudautosomal region, segment 1/2	
8681 21681	34829	1.73	1.4E+00	R20469.1	EST_HUMAN	Y933H12.11 Seires infant brain N1B Homo sapiens cDNA clone IMAGE:34345_5'	
9097 21785	34951	4.65	1.4E+00	BE084687.1	EST_HUMAN	RC1-BT013-301298-012-f05 B T013 Homo sapiens cDNA	
9131 21819	34985	0.61	1.4E+00	AF134844.1	NT	Scleropus undulatus ornithine transcarbamylase (OTC) mRNA, complete cds	

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## Single Exon Probes Expressed In Brain

Probe Seq ID No:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
10108	227556	35898	0.79	1.4E+00	BF576545.1	EST_HUMAN	602133135F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4288137 5'
10151	227899	36015	0.61	1.4E+00	BE145374.1	EST_HUMAN	IL5-HT0188-291099-008-C04 HT0188 Homo sapiens cDNA
10151	227899	36016	0.61	1.4E+00	BE145374.1	EST_HUMAN	IL5-HT0188-291099-008-C04 HT0188 Homo sapiens cDNA
10424	23070	36281	1.06	1.4E+00	D63441.1	NT	Pandorina coeruleomaculata chloroplast rbcL gene for ribulose bisphosphate carboxylase, partial cds
10424	23070	36292	1.08	1.4E+00	D63441.1	NT	Pandorina coeruleomaculata chloroplast rbcL gene for ribulose bisphosphate carboxylase, partial cds
11003	23675	36891	1.34	1.4E+00	AA195528.1	EST_HUMAN	Z33609.r1 Scores_NihMPU_S1 Homo sapiens cDNA clone IMAGE:665512 5' similar to contains element MER22 repetitive element;
11188	23833	37139	6.16	1.4E+00	AB008682.1	NT	(Homo sapiens APECED mRNA for AIRE-1, complete cds)
11381	23988	37288	4.42	1.4E+00	BE982107.2	EST_HUMAN	601655184R1 NIH_MGC_85 Homo sapiens cDNA clone IMAGE:3845805 3'
11381	23988	37289	4.42	1.4E+00	BE982107.2	EST_HUMAN	601655184R1 NIH_MGC_85 Homo sapiens cDNA clone IMAGE:3845805 3'
11404	24053	37357	3.48	1.4E+00	U30780.1	NT	Pneumocystis carinii f. sp. ratii guanine nucleotide binding protein alpha subunit (pgc1) gene, complete cds
11404	24053	37358	3.46	1.4E+00	U30780.1	NT	Pneumocystis carinii f. sp. ratii guanine nucleotide binding protein alpha subunit (pgc1) gene, complete cds
12079	25258		1.48	1.4E+00	AL161500.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 12
657	13239		1.81	1.3E+00	Z73840.1	NT	M.mucedo gene encoding 4-Dihydronathydrisporate dehydrogenase
882	13651	26320	3.42	1.3E+00	AJ271192.1	NT	Cantharellus sp. partial 2SS rRNA gene, isolate Tibet
1107	13864		20.28	1.3E+00	Y18213.1	NT	Homo sapiens putative polII HbA pseudogene for hair keratin, exons 2 to 7
1274	14024	26692	13.71	1.3E+00	4807888	NT	Homo sapiens zinc finger protein 157 (HZF22) (ZNF157) mRNA
1274	14024	26693	13.71	1.3E+00	4807888	NT	Homo sapiens zinc finger protein 157 (HZF22) (ZNF157) mRNA
1334	14083		1.28	1.3E+00	U61730.2	NT	Cox2 lacryme-pobi dihydrodipicolinate synthase (dapa) gene, complete cds
1605	14351		2.27	1.3E+00	AE00238.2	NT	Chlamydia muridarum, section 68 of the complete genome
2239	14957		1	1.3E+00	AB03047.1	NT	Cyprinus carpio MRPb and MASPB genes for mannose-binding lectin-associated serine protease (MASP)
2405	15126	27882	1.27	1.3E+00	P252301	SWISSPROT	LAMININ ALPHA-1 CHAIN PRECURSOR (LAMININ A-CHAIN)
2553	16268		1.75	1.3E+00	BE866756.2	EST_HUMAN	601661233R1 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:3916945 3'
2940	15705	28384	0.73	1.3E+00	0755621	NT	Mus musculus alpha-spectrin 1, erythroid (Spiral) mRNA
							Fugu rubripes gamma-aminobutyric acid receptor beta subunit gene, partial cds; 55kd erythrocyte membrane protein (P55), synaptic vesicle-associated integral membrane protein (VAMP-1), procollagen C-proteinase enhancer protein (PCOLCE) genes, complete cds
3534	16339	28694	0.89	1.3E+00	AF016494.1	NT	PHENOL HYDROXYLASE P3 PROTEIN (PHENOL 2-MONOXYGENASE P3 COMPONENT)
5427	18228	30938	1.09	1.3E+00	P19732	NT	Human estradiol 17 beta-dehydrogenase gene, complete cds
6522	18418	31330	0.6	1.3E+00	M27138.1	NT	602145264F1 NIH_MGC_48 Homo sapiens cDNA clone IMAGE:4309085 5'
6863	18650	31690	0.81	1.3E+00	BF663825.1	EST_HUMAN	PMD-CT0289-281188-004-108 CT0289 Homo sapiens cDNA
6928	18712	31687	7.57	1.3E+00	AW3622834.1	EST_HUMAN	PMD-CT0289-281188-004-108 CT0289 Homo sapiens cDNA

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## Single Exon Probes Expressed In Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
5923	18712	31698	7.57	1.3E+00	AW362834.1	EST_HUMAN	P00-CT0289-291189-004-f08 CT0289 Homo sapiens cDNA D.melanogaster no-on-transient A gene product, complete cds
6323	19093	32081	1.34	1.3E+00	MS3496.1	NT	
6682	18414		0.76	1.3E+00	Q00156	SWISSPROT	HYPOTHETICAL PROTEIN
6738	18573	32606	0.62	1.3E+00	M13918.2	NT	Homo sapiens fibronectin receptor, alpha-subunit precursor (ITGA5) mRNA, partial cds
6884	19554	32584	1.17	1.3E+00	BE638819.1	EST_HUMAN	601061420F1 NIH_MGSC_10 Homo sapiens cDNA clone IMAGE:3447885 5'
7000	18892	32743	0.81	1.3E+00	BE243571.1	EST_HUMAN	TCBAP1D0859 Pediatric pre-B cell acute lymphoblastic leukemia Baylor-HGSC project=TCBA Homo sapiens cDNA clone TCBAP1D0859
7358	20039	33117	1.01	1.3E+00	P24540	SWISSPROT	ACYLPHOSPHATASE, ORGAN-COMMON TYPE ISOZYMES A AND B (ACYLPHOSPHATE PHOSPHOHYDROLASE)
8197	20891	34029	1.23	1.3E+00	AJ008912.1	NT	Sus scrofa pip gene
8348	21039	34178	2.78	1.3E+00	BE983378.2	EST_HUMAN	601657145R1 NIH_MGSC_67 Homo sapiens cDNA clone IMAGE:38681165 3'
8459	21151	34294	0.88	1.3E+00	BE974280.1	EST_HUMAN	601680250R2 NIH_MGSC_83 Homo sapiens cDNA clone IMAGE:3950532 3'
8811	21303		1.78	1.3E+00	8910247	NT	Homo sapiens GL004 protein (GL004), mRNA
8889	21381	34525	0.78	1.3E+00	A1827629.1	EST_HUMAN	W085a073X1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2482100 3'
9415	22093		5.24	1.3E+00	AF042084.1	NT	Homo sapiens heparan glucosaminyl N-deacetylase/N-sulfotransferease-2 gene, complete cds
9424	22102	35273	2.68	1.3E+00	X72018.1	NT	S.alba phr-1 mRNA for photophase
9424	22102	35274	2.58	1.3E+00	X72018.1	NT	Homo sapiens lipoygenase (ALOX12B) mRNA, complete cds
9524	22117	35381	0.98	1.3E+00	AF059250.1	NT	S.alba phr-1 mRNA for photophase
9589	22222	35407	1.58	1.3E+00	O00754	SWISSPROT	LYSOSOMAL ALPHA-MANNOSEIDASE PRECURSOR (MANNOSIDASE, ALPHA B) (LYSOSOMAL ACID ALPHA-MANNOSEIDASE) (LAMAN)
9651	22303	35498	1.14	1.3E+00	AI827628.1	EST_HUMAN	W085a073X1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2482100 3'
9726	22377	35578	0.79	1.3E+00	AJ223982.1	NT	Lactococcus lactis cremoris NCDO-lvrl chromosome I inversion junction DNA
9726	22377	35579	0.79	1.3E+00	AJ223982.1	NT	Lactococcus lactis cremoris NCDO-lvrl chromosome I inversion junction DNA
9766	22447	35624	4.53	1.3E+00	BE863378.2	EST_HUMAN	601657145R1 NIH_MGSC_67 Homo sapiens cDNA clone IMAGE:38681165 3'
9826	22477		0.48	1.3E+00	AJ569844.1	EST_HUMAN	W0774d2X1 NCI_CGAP_UH Homo sapiens cDNA clone IMAGE:2214814 3' similar to gb:X14723 CLUSTERIN PRECURSOR (HUMAN)
100050	22698	35913	0.46	1.3E+00	AF081251.1	NT	Escherichia coli serotype O157:H7 O antigen gene cluster
10050	22698	35914	0.46	1.3E+00	AF081251.1	NT	Escherichia coli serotype O157:H7 O antigen gene cluster
10113	22761	35974	1.62	1.3E+00	AE004392.1	NT	Vibrio cholerae chromosome I, section 49 of 83 of the complete chromosome
10130	22778	35981	1.35	1.3E+00	M28953.1	NT	Campylobacter jejuni karamayica phosphotransferase (alpha-7) gene, complete cds
10483	23128		0.82	1.3E+00	AL16533022	NT	Homo sapiens chromosome 21 segment HS2C102
10511	23157	36383	0.45	1.3E+00	AI890846.1	EST_HUMAN	W032e10-X1 NCI_CGAP_GCG Homo sapiens cDNA clone IMAGE:2498922 3' similar to SW:TRXB_HUMAN Q16881 THIOPEROXIDIN REDUCTASE
10592	23286		4.6	1.3E+00	Q14117	SWISSPROT	DHYDROXYRIMIDINASE (DHPase) (HYDANTOINASE) (DHP)

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**Table 4**  
**Single Exon Probes Expressed In Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
10822	23505	36744	1.93	1.3E+00	P25289	SWISSPROT	mRNA 3'-END PROCESSING PROTEIN RNA15
10846	23523	36772	2.01	1.3E+00	Z18892.2	NT	Mus musculus desmin gene
11307	23866		1.8	1.3E+00	AW274791.1	EST_HUMAN	xp09503_x1 NCL CGAP_HNB Homo sapiens cDNA clone IMAGE:2739868 3'
11527	24127	37433	3.21	1.3E+00	D42042.1	NT	Human mRNA for KIAA0085 gene, partial cds
11624	24221	37544	3.18	1.3E+00	Z98682.1	NT	Bacillus subtilis genomic DNA 23_9kB fragment
12210	24675		2.64	1.3E+00	AF187873.1	NT	Cavia porcellus inwardly-rectifying potassium channel Kir2.2 (KCNU12) gene, complete cds
12386	24730	31035	0.3	1.3E+00	BF348043.1	EST_HUMAN	602023185F1 NCI CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4158452 5'
12397	25153		2.73	1.3E+00	F33484	SWISSPROT	E1 GLYCOPROTEIN PRECURSOR (MATRIX GLYCOPROTEIN) (MEMBRANE GLYCOPROTEIN)
12489	24848		2.15	1.3E+00	AF187035.1	NT	Stomaria lilium cytochrome b gene, complete cds; mitochondrial gene for mitochondrial product
635	13444	26050	11.05	1.2E+00	AA876248.1	EST_HUMAN	212208.s1 Soares, fetal liver spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:4316363 3'
804	13576	26239	0.87	1.2E+00	PO5228	SWISSPROT	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-II)
804	13576	26240	0.87	1.2E+00	PO5228	SWISSPROT	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-III)
804	13576	26241	0.87	1.2E+00	PO5228	SWISSPROT	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-III)
858	13627		1.35	1.2E+00	8924234	NT	Homo sapiens hypothetical protein PRO3077 (PRO3077), mRNA
1138	13883	26554	5.64	1.2E+00	AF080245.2	NT	Eleotris difesa sesquiterpene synthase mRNA, complete cds
1183	13985	26890	1.28	1.2E+00	AJ252242.1	NT	pea seed-borne mosaic virus complete genome
1183	13985	26891	1.28	1.2E+00	AJ252242.1	NT	pea seed-borne mosaic virus complete genome
2003	14739	27483	1.22	1.2E+00	AF140631.1	NT	Homo sapiens G-protein coupled receptor 14 (GPR14) gene, complete cds
9108	15873	28512	1.24	1.2E+00	AB020881.1	NT	Homo sapiens mRNA for KIAA0874 protein, partial cds
3163	15926	28573	5.88	1.2E+00	AL161563.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 63
3163	15926	28574	5.88	1.2E+00	AL161563.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 63
3280	16041		2.59	1.2E+00	P54890	SWISSPROT	CONJUGAL TRANSFER PROTEIN TRIBE PRECURSOR
3689	18452	28091	6.69	1.2E+00	U75902.1	NT	Mus musculus subtilisin-like serine protease LPC (PC7) gene, exons 1 to 9, partial cds
3987	18716	28354	1.78	1.2E+00	BF373570.1	EST_HUMAN	MRO-FT075-050900-203_906_1 FT0175 Homo sapiens cDNA
4266	16110	28783	1.11	1.2E+00	AF188740.1	NT	Homo sapiens Lhx3 gene, intron 2
4438	17174		1.67	1.2E+00	M87080.1	NT	Rattus rattus cardiac Ace3 gene, exons 1-23
4487	17222	28850	0.86	1.2E+00	AL161509.2	NT	Arachidopsis italiana DNA chromosome 4, contig fragment No. 21
4523	17258	28892	1.89	1.2E+00	AF168495.1	NT	Homo sapiens post-synaptic density 95 (DLG4) gene, complete cds
4548	17283		0.43	1.2E+00	Y09200.1	NT	T-plastidium chloroplast rbcL gene, partial
5351	18154	30836	1.1	1.2E+00	U20760.1	NT	Human extracellular calcium-sensing receptor mRNA, complete cds
5487	18268	31158	1.91	1.2E+00	AW813276.1	EST_HUMAN	MR3-ST0181-1-40200-013-005 S70191 Homo sapiens cDNA
5784	18576	31504	0.83	1.2E+00	AF016052.1	NT	Homo sapiens zinc finger protein ZNF191 (ZNF191) gene, complete cds
6060	18840	31801	2.51	1.2E+00	X74885.1	NT	D. hydei avl repeat cluster DNA, fragment D
6119	18897	31885	4.42	1.2E+00	BE003113.1	EST_HUMAN	QV4-BN0090-27040-190-403 BN0090 Homo sapiens cDNA

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Table 4  
Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
6198 18974		31951	1.54	1.2E+00	X89084.1	NT	C. glutamicum pba gene and ackA gene
6199 18974		31952	1.54	1.2E+00	X89084.1	NT	C. glutamicum pba gene and ackA gene
6241 19015		31969	39.54	1.2E+00	AA759254.1	EST_HUMAN	ah84g12.s1 Soares testis NHT Homo sapiens cDNA clone 13223743'
6342 19112		32101	0.55	1.2E+00	N39225.1	EST_HUMAN	Y38b12.s1 Soares fibroblastocyte 2NIHM Homo sapiens cDNA clone IMAGE:273598 3' similar to gb M87935 HUMAU472 Human carcinoma cell-derived Alu RNA transcript, (rRNA); gb J04970 CARBOXYPEPTIDASE M PRECURSOR (HUMAN);
6408 19177		32175	0.88	1.2E+00	P17671	SWISSPROT	ECDSONE-INDUCIBLE PROTEIN E75-A
6412 19180		32179	2.06	1.2E+00	AW813276.1	EST_HUMAN	MR2-ST0191-140200-013-005 ST0191 Homo sapiens cDNA
6815 19376		32498	1.17	1.2E+00	AB029010.1	NT	Homo sapiens mRNA for KIAA1087 protein, partial cds
6826 19490		32512	3.11	1.2E+00	AJ002141.1	NT	Mus musculus DSPP gene
7163 19840			0.84	1.2E+00	AJ271795.1	NT	Homo sapiens Xq pseudautosomal region; segment 1/2
7282 25109		33044	4.88	1.2E+00	AV734585.1	EST_HUMAN	AV734585 cDNA Homo sapiens cDNA clone cAA1F-035 5'
7650 20220		33323	2.49	1.2E+00	X74207.1	NT	Lactis pyD and pyF genes
7603 20269		33376	0.58	1.2E+00	J05218.1	NT	Chicken muscarinic acetylcholine receptor (cm4 MACHR) gene, complete cds
7715 20379		33492	0.58	1.2E+00	BET878784.1	EST_HUMAN	601487761F1 NIH MGIC 88 Homo sapiens cDNA clone IMAGE:3884270 5'
8467 21159		34302	3.32	1.2E+00	AB033030.1	NT	Homo sapiens mRNA for KIAA1204 protein, partial cds
8551 21253		34391	0.68	1.2E+00	P38427	SWISSPROT	ALPHA-ALPHA-TREHALOSE-PHOSPHATE SYNTHASE [UDP-FORMING] 123 KD SUBUNIT (TREHALOSE-6-PHOSPHATE SYNTHASE) (UDP-GLUCOSE-GLUCOSEPHOSPHATE GLUCOSYLTRANSFERASE)
8776 21497			0.51	1.2E+00	7708271	NT	Homo sapiens Cgl-30 protein (LOC51811), mRNA
8923 21614		34768	1.87	1.2E+00	AW377210.1	EST_HUMAN	MR2-CT0222-201088-01-07 CT0222 Homo sapiens cDNA
9138 21626		34991	0.5	1.2E+00	H48389.1	EST_HUMAN	Y580a08.1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:202068 6
9228 21985		35138	3.76	1.2E+00	Z32850.1	NT	R.commuins gene for pyrophosphate-dependent phosphofructokinase beta subunit
9505 22158		35339	1.81	1.2E+00	D11745.1	EST_HUMAN	HUM-MMO1A01 Liver HepG2 cell line, Homo sapiens cDNA clone hm01a01
9831 22482		35684	2.88	1.2E+00	Y568632.1	NT	H.sapiens ENO3 gene for muscle specific endopeptidase
10224 22872			0.73	1.2E+00	AB008886.1	NT	Homo sapiens Klotho gene, exon 1
11318 24009		37314	3.78	1.2E+00	AW817817.1	EST_HUMAN	PM0-ST0284-161188-001-01 ST0284 Homo sapiens cDNA
11357 24045			10.82	1.2E+00	BE160781.1	EST_HUMAN	PM1-HT0422-160200-007-810 HT0422 Homo sapiens cDNA
11435 28202		36434	4.36	1.2E+00	U50147.1	NT	Retius norvegicus synapse-associated protein 102 mRNA, complete cds
12179 25227		30817	17.06	1.2E+00	AL163203.2	NT	Homo sapiens chromosome 21 segment HS21C003
12189 24687			2.8	1.2E+00	AP001515.1	NT	Bacillus halodurans genomic DNA, section 9/14
451 13237		25876	1.53	1.1E+00	D88980.1	NT	Human mRNA for KIAA0227 gene, partial cds
1767 14499		27200	1.33	1.1E+00	AW985383.1	EST_HUMAN	QVO-BN0042-170300-163-912 BN0042 Homo sapiens cDNA
1892 14629		27339	0.98	1.1E+00	AW575888.1	EST_HUMAN	UHF-BR0-pkk-f02-qJL.st1 NIH MGIC 52 Homo sapiens cDNA clone IMAGE:3074834 3'

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Table 4 Single Event Sensitive

Single Locus Probes Expressed in Brain							
Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
3324	16084	28734	6.48	1.1E+00	AL163213.2	NT	Homo sapiens chromosome 21 segment HS21C013
3324	16084	28735	6.48	1.1E+00	AL163213.2	NT	Homo sapiens chromosome 21 segment HS21C013
3480	16236	28892	1.11	1.1E+00	8922841	NT	Homo sapiens hypothetical protein FLJ10749 (FLJ10749), mRNA
3587	16322	28970	1.01	1.1E+00	AI808360.1	EST_HUMAN	wf54h11.x1 Scores_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2359461 3' similar to SW:P531_HUMAN Q12888 PG3-BINDING PROTEIN 33P1 ;
3707	16460	28098	1.05	1.1E+00	AE038886.1	NT	Xylella fastidiosa, section 32 of 229 of the complete genome
3707	16460	20099	1.05	1.1E+00	AE038886.1	NT	Xylella fastidiosa, section 32 of 229 of the complete genome
3768	16550		1.02	1.1E+00	XB3574.1	NT	H.paraaeruginosa phbII(C), hphII(C), hphII(R) and manB genes
4190	16831		5.68	1.1E+00	68363331	NT	R.unimoriis complete mitochondrial genome
4634	17369		0.81	1.1E+00	U34862.1	NT	Cancherhuius plumbeus Ig lambda light chain gene, complete cds
4934	17682	30272	3.45	1.1E+00	U18486.1	NT	African swine fever virus, complete genome
4935	17683	30273	1.05	1.1E+00	AJ271740.1	NT	Drosophila melanogaster D-Titin gene, exons 1-37
5128	17847	30484	1.07	1.1E+00	68890086	NT	Homo sapiens putative GR6 protein (GR6), mRNA
5224	18031	30657	1.39	1.1E+00	69785330	NT	Rattus norvegicus Aquaporin 4 (Aqp4), mRNA
6528	18324	31226	15.75	1.1E+00	BE80184.1	EST_HUMAN	6018652776R1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:3826835 3'
6545	18342	31260	1.2	1.1E+00	AI136882.1	EST_HUMAN	qd85c03.x1 Scores_tests_NHT_Homo sapiens cDNA clone IMAGE:7362650 3'
6601	18782	31743	1.1	1.1E+00	11419739	NT	Homo sapiens solute carrier family 6 (neurotransmitter transporter), member 14 (SLC6A14), mRNA
6181	18958	31932	0.62	1.1E+00	AF197881.1	NT	Macgregoria pulchra cytochrome b gene, complete cds; mitochondrial gene for mitochondrial product yeast03.c1 Scores fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:124924 5'
6313	19084	32069	0.82	1.1E+00	RG6037.1	EST_HUMAN	Mus musculus mRNA for ER protein 58 (EP58 gene)
6616	19379	32294	0.72	1.1E+00	AI404004.1	NT	Homo sapiens collagen type XI alpha-1 (COL11A1) gene, exons 25 through 28
7155	19842		0.58	1.1E+00	AF010691.1	NT	Maize mRNA for endopeptidase (2'-phospho-D-glucosidase hydrolase)
7196	19882	32958	0.72	1.1E+00	X55881.1	NT	Herpes simplex virus type 1 (strain KOS) UL41 gene
7389	20168	33146	2.18	1.1E+00	272398.1	NT	Herpes simplex virus type 1 (strain KOS) UL41 gene
7389	20068	33147	2.18	1.1E+00	272398.1	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 84
7411	20088	33172	8.84	1.1E+00	AL161988.2	NT	Mus musculus silent mating type information regulation 2, (S.cerevisiae, homolog)-like (SI22), mRNA
7480	25115	33247	0.8	1.1E+00	11987980	NT	602082562F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4248828 5'
8032	20727	33860	3.01	1.1E+00	BF688986.1	EST_HUMAN	IntS11.X1 NCI_CGAP_Kdr1 Homo sapiens cDNA clone IMAGE:2160349 3'
8120	20814	33950	0.84	1.1E+00	AI78339.1	EST_HUMAN	Acetabularia caliculus mitochondrial COX1-like gene
8638	21328	34471	0.71	1.1E+00	AB030088.1	NT	VH=anti-cytomegalovirus glycoprotein B antibody 4D4 heavy chain variable region [Human, mRNA Partial], 375 nt]
8714	21408	34549	0.75	1.1E+00	S80750.1	NT	

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Table 4  
Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
8824 21516	34681	0.45	1.1E+00	AJ079846.1	EST_HUMAN	023405_x1 Soares_NIHMPu_S1 Homo sapiens cDNA clone IMAGE:1677249 3'	
8837 20408		0.69	1.1E+00	BE384878.1	EST_HUMAN	6012176278f1 NIH_MGC_20 Homo sapiens cDNA clones IMAGE:3617418 5'	
8828 22181	35365	0.53	1.1E+00	AJ245772.1	NT	Mus musculus mRNA for stretch responsive muscle (X-chromosome) protein (Smox gene)	
98580 22233		1.2	1.1E+00	Y12227.1	NT	Arabidopsis thaliana DNA, 24 kb surrounding PFL locus	
9872 22324	36520	1.14	1.1E+00	L78301.1	NT	Yersinia pseudotuberculosis psaE, psaF, usherin (psaB), chaperone (psaA), and usher (psaC) genes, complete cds	
9732 22383	35585	1.37	1.1E+00	AB023151.1	NT	Homo sapiens mRNA for KIAA0834 protein, partial cds	
8837 22488	35690	4.59	1.1E+00	AL161516.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 27	
9898 22548	35742	18.34	1.1E+00	6754021	NT	Mus musculus glutamine nucleotide binding protein (G protein), gamma 3 subunit (Gng3), mRNA	
10398 23044	36260	1.1	1.1E+00	P73769	SWISSPROT	DNase mismatch repair protein MUTS	
10504 23150	36375	0.73	1.1E+00	AI878821.1	EST_HUMAN	as5ic1.1_Y Schneider fetal brain 00004 Homo sapiens cDNA clone IMAGE:2518292 & similar to gb:D10522 Human mRNA for 80K-L protein, complete cds. (HUMAN)	
10547 23243	36478	2.25	1.1E+00	11067384	NT	Homo sapiens KIAA0828 gene product (KIAA0828), mRNA	
10606 23300		3.1	1.1E+00	AF088942.1	NT	Klebsiella pneumoniae flavins cytochrome c oxidase subunit 2 (cox2) gene, mitochondrial gene encoding mitochondrial protein, partial cds	
11023 23695	36958	1.28	1.1E+00	11439586	NT	Homo sapiens potassium inwardly-rectifying channel, subfamily J, member 11 (KCNJ11), mRNA	
11025 23698	36981	1.58	1.1E+00	L16877.1	NT	Homo sapiens cytochrome P4502C9 (CYP2C9) gene, 5' flanking and exon 1	
11042 17901		6.23	1.1E+00	8922873	NT	Homo sapiens hypothetical protein FLJ11280 (FLJ11280), mRNA	
11048 23718	36988	3.88	1.1E+00	AF012862.1	NT	Petroselinum crispum cytosolic glucose-6-phosphate dehydrogenase 1 (cG6PDH1), mRNA, complete cds	
11048 23718	36989	3.88	1.1E+00	AF012862.1	NT	Petroselinum crispum cytosolic glucose-6-phosphate dehydrogenase 1 (cG6PDH1) mRNA, complete cds	
11328 24019	37323	4.58	1.1E+00	AI808898.1	EST_HUMAN	w7f611x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2361548 3'	
11561 24160	37470	1.63	1.1E+00	D89501.1	NT	Human PR1 gene, complete cds	
11561 24160	37471	1.63	1.1E+00	D89501.1	NT	Human PR1 gene, complete cds	
12153 24639		3.88	1.1E+00	P07886	SWISSPROT	LOW TEMPERATURE ESSENTIAL PROTEIN	
12250 24697	31078	1.93	1.1E+00	AF216898.1	NT	Taenia solium immunogenic protein Ts76 mRNA, partial cds	
12378 25225		2.09	1.1E+00	AF234469.1	NT	Dichotomius discoidalis isopenetyl pyrophosphate isomerase (Dipi) mRNA, complete cds	
12388 25200		1.44	1.1E+00	8393198	NT	Rattus norvegicus C-reactive protein, member of the pentraxin family (Crp), mRNA	
97 12923		2.48	1.0E+00	U233808.1	NT	Xenopus laevis modopin gene, complete cds	
111 12852	25569	0.73	1.0E+00	D88425.1	NT	Cavia cobaya mRNA for serine/threonine kinase, complete cds	
409 13184		2.25	1.0E+00	AB021684.1	NT	Marchantia polymorpha genes for 28S rRNA, 18S rRNA, 6.8S rRNA and 26S rRNA	
562 13344	25971	1.2	1.0E+00	AJ251680.1	NT	Giardia lamblia mRNA for homeodomain transcription factor (so gene)	
682 13458	26079	4.38	1.0E+00	AL163218.2	NT	Homo sapiens chromosome 21 segment HS21C018	

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Table 4  
Single Exon Probes Expressed in Brain

Probe Seq ID No:	Exon Seq ID No:	ORF Seq ID No:	Expression Signal	Most Similar BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
663	13439		0.95	1.0E+00 AF25984.1	NT	Aedes aegypti mucin-like protein MUC1 mRNA, complete cds	
1365	15567		3.03	1.0E+00 X80416.1	NT	V.vulgaris Agar-CAM mRNA	
1751	14463	27183	0.83	1.0E+00 AB008531.1	NT	<i>Platirus stellatus</i> intestine virus RNA for nonstructural polyprotein, capsid protein precursor, complete cds	
2489	16208	27847	1.18	1.0E+00 P48355	SWISSPROT	DNA GYrase subunit B	
2489	15208	27848	1.18	1.0E+00 P48355	SWISSPROT	DNA GYrase subunit B	
2878	15645	28287	3.82	1.0E+00 P24008	SWISSPROT	3-OXO-5-ALPHA-STEROID 4-DEHYDROGENASE 1 (STEROID 5-ALPHA-REDUCTASE 1) (SR TYPE 1)	
2878	16845	28288	3.82	1.0E+00 P24008	SWISSPROT	3-OXO-5-ALPHA-STEROID 4-DEHYDROGENASE 1 (STEROID 5-ALPHA-REDUCTASE 1) (SR TYPE 1)	
2967	15733		1.17	1.0E+00 O14228	SWISSPROT	HYPOTHETICAL 67.9 KD PROTEIN C6F12.08C IN CHROMOSOME 1	
3194	15957	26609	1.24	1.0E+00 AA622453.1	EST_HUMAN	ef52g08_s1 Scores_total_fatus_Nb2HFB_BW Homo sapiens cDNA clone IMAGE:1032830 3' similar to WP:CA2D8.3 CE04204: contains element MER22/MER22 repetitive element;	
3685	12923		1.24	1.0E+00 U23808.1	NT	Xenopus laevis rhodopsin gene, complete cds	
3689	16422	28063	1.04	1.0E+00 AJ223816.1	NT	<i>Agelaius phoeniceus</i> mRNA for tyrosinase	
4050	16795	28424	0.76	1.0E+00 AF223391.1	NT	Homo sapiens calcium channel alpha1E subunit (CACNA1E) gene, exons 7-49, and partial cds, alternatively spliced	
4242	16983		0.79	1.0E+00 68222245	NT	Homo sapiens hypothetical protein FLJ10139 (FLJ10139), mRNA	
4954	17680		0.83	1.0E+00 D10852.1	NT	Rattus norvegicus mRNA for N-acetylglucosaminyltransferase II, complete cds	
4975	17698	30303	0.74	1.0E+00 AF092505.1	NT	Mus musculus dipeptidyl aminopeptidase-like protein 6 (Dpp6) gene, partial cds; and prostatic Rump white inversion breakpoint	
5220	18008	30629	3.63	1.0E+00 Z97022.1	NT	<i>Hordeum vulgare</i> gene encoding cysteine proteinase	
5759	18551	31472	4.97	1.0E+00 AF248054.1	NT	Bos taurus microtumor calcium activated neutral protease 1 (CAPN1) gene, exons 11-20, and partial cds	
5759	18551	31473	4.97	1.0E+00 AF248054.1	NT	Bos taurus microtumor calcium activated neutral protease 1 (CAPN1) gene, exons 11-20, and partial cds	
5897	18654	31695	1.53	1.0E+00 297341.2	NT	<i>Arabidopsis thaliana</i> DNA chromosome 4, ESSA 1 FCA contig fragment No. 6	
6024	18804	31765	4.7	1.0E+00 P04501	SWISSPROT	FIBER PROTEIN	
6030	18810	31770	1.49	1.0E+00 AW452782.1	EST_HUMAN	U1-H-B13-alk-d-09-04/J_91 NCI CGAP_Subs Homo sapiens cDNA clone IMAGE:30188968 3'	
6397	18168	32168	1.95	1.0E+00 U75802.1	NT	Mus musculus subtilisin-like serine protease LPC (PCT) gene, exons 1 to 6, partial cds	
6447	18215	32213	0.91	1.0E+00 AF104882.1	NT	<i>Homo sapiens</i> cell cycle protein (PA2G4) gene, exons 2 through 5	
6534	18300		1.08	1.0E+00 P46508	SWISSPROT	SRB-11 PROTEIN	
6579	18598	32634	1.33	1.0E+00 Y11204.1	NT	V.vulgaris gene encoding vacuole	

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**Table 4**  
**Single Exon Probes Expressed in Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
7038	19730	32788	1.09	1.0E+00	SS2770.1	NT	Inslin-like growth factor-binding protein 4 [cattle, pulmonary artery endothelial cells, mRNA, 2028 nt]
7378	20058		8.29	1.0E+00	F20273	SWISSPROT	B-CELL RECEPTOR CD22 PRECURSOR (LEU-14) (B-LYMPHOCYTE CELL ADHESION MOLECULE) (BL-CAM)
7811	20277	33385	1.36	1.0E+00	AF192331.1	NT	Homo sapiens endothelin-converting enzyme 2 (ECE2) mRNA, complete cds
7628	20292	33401	6.26	1.0E+00	AA775181.1	EST_HUMAN	ac79608.31 Strategene lung (#337210) Homo sapiens cDNA clone IMAGE:888791 3'
7881	20558	33681	1.36	1.0E+00	BE868267.1	EST_HUMAN	601443850F1 NIH MGIC_65 Homo sapiens cDNA clone IMAGE:3848006 5'
7881	20558	33682	1.36	1.0E+00	BE868267.1	EST_HUMAN	601443850F1 NIH MGIC_65 Homo sapiens cDNA clone IMAGE:3848005 5'
8041	17880		1.19	1.0E+00	D10852.1	NT	Rattus norvegicus mRNA for N-acetylglucosaminyltransferase III, complete cds
8248	20942	34079	2.02	1.0E+00	Q02207	SWISSPROT	PEROXISOMAL HYDRATASE-DEHYDROGENASE-EPIMERASE (HDE) (MULTIFUNCTIONAL BETA-OXIDATION PROTEIN) (MFP) [INCLUDES: 2-ENOYL-COA HYDRATASE ; D-3-HYDROXYACYL COA DEHYDROGENASE ]
8248	20942	34080	2.02	1.0E+00	Q02207	SWISSPROT	PEROXISOMAL HYDRATASE-DEHYDROGENASE (HDE) (MULTIFUNCTIONAL BETA-OXIDATION PROTEIN) (MFP) [INCLUDES: 2-ENOYL-COA HYDRATASE ; D-3-HYDROXYACYL COA DEHYDROGENASE ]
8378	21089		0.85	1.0E+00	F51784	SWISSPROT	UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 11 (UBIQUITIN THIOLESTERASE 11) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 11) (DEUBIQUITINATING ENZYME 11)
8408	21101	34237	0.5	1.0E+00	C9Y5T5	SWISSPROT	UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 16 (UBIQUITIN THIOLESTERASE 16) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 16) (DEUBIQUITINATING ENZYME 16) (UBIQUITIN PROCESSING PROTEASE UBP-M)
8408	21101	34238	0.5	1.0E+00	O8Y6T6	SWISSPROT	UBIQUITIN CARBOXYL-TERMINAL HYDROLASE 18 (UBIQUITIN THIOLESTERASE 18) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE 18) (DEUBIQUITINATING ENZYME 18) (UBIQUITIN PROCESSING PROTEASE UBP-M)
8436	26122		2.34	1.0E+00	BE147331.1	EST_HUMAN	RC1-HT0229-181089-011-003 HT0229 Homo sapiens cDNA
8478	21168	34312	0.88	1.0E+00	U42720.2	NT	Simian immunodeficiency virus Gag protein (gag) gene, complete cds; Poi protein (poi) gene, partial cds; and Vif protein (vif), Vpr protein (vpr), Tat protein (tat), Rev protein (rev), Vpu protein (vpu), Env protein (env), and Nef protein (nef) genes. >
8625	21317	34459	1.27	1.0E+00	M38427.1	EST_HUMAN	Human immunodeficiency virus type 1 (HIV-1), Isolates SF233,
9171	21841	35003	2.43	1.0E+00	BE907592.1	EST_HUMAN	601497581F1 NIH MGIC_70 Homo sapiens cDNA clone IMAGE:3889021 5'
9381	22043	36213	1.69	1.0E+00	6753429	NT	Mus musculus chloride channel calcium activated 1 (Clca1), mRNA
9381	22043	35218	1.69	1.0E+00	6753429	NT	Mus musculus chloride channel calcium activated 1 (Clca1), mRNA
9510	22163	35345	1.83	1.0E+00	AV6889554.1	EST_HUMAN	AV6889554 GKC Homo sapiens cDNA clone GKCCYCA15
9516	22168	35351	1.43	1.0E+00	U44952.1	NT	Xenopus laevis zona pellucida C glycoprotein precursor (xZPC) mRNA, complete cds
9516	22168	35352	1.43	1.0E+00	U44952.1	NT	Xenopus laevis zona pellucida C glycoprotein precursor (xZPC) mRNA, complete cds

**Table 4**  
**Single Exon Probes Expressed in Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
9753	22404	35608	0.49	1.0E+00	X15498.1	NT	Human Coronavirus gene for membrane protein
9753	22404	35610	0.49	1.0E+00	X15498.1	NT	Human Coronavirus gene for membrane protein
10012	22860	35875	0.71	1.0E+00	5174562	NT	Homo sapiens MHC binding factor, beta (MHCBFB) mRNA
10012	22860	35876	0.71	1.0E+00	5174562	NT	Homo sapiens MHC binding factor, beta (MHCBFB) mRNA
10104	22752	35988	0.81	1.0E+00	A107720.1	EST_HUMAN	c916d07.s1 Soares_scenescent_fibroblasts_NBHSF Homo sapiens cDNA clone IMAGE:16858013
10226	22873	36085	4.36	1.0E+00	AV7558825.1	EST_HUMAN	AV7558825 BM1 Homo sapiens cDNA clone BMFFAWC04 6'
10375	23021	36237	16.16	1.0E+00	AA004982.1	EST_HUMAN	ZB94e02.11 Soares_fetal_liver_spleen_1NF1.S1 Homo sapiens cDNA clone IMAGE:4228908 5'
10375	23021	36238	16.16	1.0E+00	AA004982.1	EST_HUMAN	ZB94e02.11 Soares_fetal_liver_spleen_1NF1.S1 Homo sapiens cDNA clone IMAGE:4228908 5'
10407	23053	36270	1.1	1.0E+00	L11810.1	NT	Human ratheoblastoma susceptibility gene exons 1-27, complete cds
10893	23573	36823	4.57	1.0E+00	S90828.1	NT	PBR17-proline-rich protein [fratton 3] [human, Geriatric, 886 n]
11025	23897	36960	1.49	1.0E+00	AA701494.1	EST_HUMAN	Z63b11.s1 Soares_fetal_liver_spleen_1NF1.S1 Homo sapiens cDNA clone IMAGE:425453 3' similar to contains Alu repetitive element; contains element MER38 repetitive element;
11522	24122		1.59	1.0E+00	L47613.1	NT	Plata glauca ElM13 mRNA
11744	18008	30639	1.55	1.0E+00	Z970722.1	NT	Hordeum vulgare gene encoding cysteine protease
11838	24422	37768	12.29	1.0E+00	Q800119	SWISSPROT	NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 8 (NADH DEHYDROGENASE 1, CHAIN 8) (NDH-1, CHAIN 8)
11868	24452	37794	1.38	1.0E+00	8826187	NT	Human adenovirus type 5, complete genome
12049	24568		3.01	1.0E+00	P153038	SWISSPROT	THROMBOMODULIN PRECURSOR (FETOMODULIN) (TM)
12370	24772		2.32	1.0E+00	AW979184.1	EST_HUMAN	EST38e283 MAGE resequences, MAGN Homo sapiens cDNA
2843	15353	28097	1.19	8.8E-01	AL163302.2	NT	Homo sapiens chromosome 21 segment HS21C102
2561	16345		0.97	9.0E-01	AF74585.1	NT	Apple mosaic virus RNA 2 putative polymerase gene, complete cds
5547	18344	31253	10.09	9.8E-01	P49857	SWISSPROT	SERINE/TREONINE PROTEIN KINASE MINIBRAIN
5778	18570	31498	0.83	9.8E-01	Q086532	SWISSPROT	PROBABLE OXIDOREDUCTASE ZK1280.5 IN CHROMOSOME II
9160	21830		1.37	9.9E-01	U656687.1	NT	Lycopersicum esculentum putative Mt1 copy 1 nematode-resistance gene
9455	22035		2.18	9.9E-01	Q288442	SWISSPROT	B2 BRADYKININ RECEPTOR (BK-2 RECEPTOR)
10614	23308	38547	2.37	9.9E-01	AJ005028.1	NT	Danio rerio mRNA for Eph-like receptor tyrosine kinase rk8
11592	24101	37508	2.3	9.9E-01	Y11972.1	NT	B.aphidicola 16S rDNA (host T. subter)
11592	24101	37509	2.3	9.9E-01	Y11972.1	NT	B.aphidicola 16S rDNA (host T. subter)
510	13294	25528	1.14	9.8E-01	P222507	SWISSPROT	AMINO ACID ACETYLTRANSFERASE (N-ACETYLGUTAMATE SYNTHASE) (AGS) (NAGS)
2285	16020		1.21	9.8E-01	AJ003108.1	NT	Callithrix jacchus UBE1 gene derived retropon on the Y chromosome
2804	16509		1.01	9.8E-01	AF174844.1	NT	Xenopus laevis rac GTPase mRNA, complete cds
3781	16533	29171	0.92	9.8E-01	O87551	SWISSPROT	PROBABLE ENDONUCLEASE IV (ENDODEOXYRIBONUCLEASE IV)
7099	19798	32852	4.67	9.8E-01	AJ302158.1	NT	Enterobacteriaceae sp. JM103 partial groES gene for GroEL-like protein, Isolate JM103

Table 4

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Meet Similar BLAST E Value	(Top) Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
7069	19788	32853	4.67	9.8E-01	AJ302158.1	NT	Enterobacteriaceae sp. JM983 partial groES gene for GroES-like protein and partial groEL gene for GroEL-like protein, Isolate JM983
7545	20215	33316	1.15	9.8E-01	BF034016.1	EST_HUMAN	601456337F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3860049 5'
7645	20216	33317	1.16	9.8E-01	BF034016.1	EST_HUMAN	601456337F1 NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3860049 5'
8619	21311	34453	0.91	9.8E-01	P38652	SWISSPROT	PHOSPHOGLUCOMUTASE (GLUCOSE PHOSPHOMUTASE) (PGM)
10336	22883		1.13	9.8E-01	AA825585.1	EST_HUMAN	cd55d04_a1 NCI_CGAP_GcB1 Homo sapiens cDNA clone IMAGE:1371847 3'
10916	23598	36842	2.29	9.8E-01	BE258705.1	EST_HUMAN	601110258F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350750 5'
10919	23599	36843	2.29	9.8E-01	BE258705.1	EST_HUMAN	601110258F1 NIH_MGC_16 Homo sapiens cDNA clone IMAGE:3350750 5'
11764	24355	37688	1.57	9.8E-01	AJ680878.1	EST_HUMAN	bx2c10x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:2272242 3'
12256	24702		1.58	9.8E-01	U52111.2	NT	Homo sapiens X28 region near ALD locus containing dual specificity phosphatase 9 (DUSP9), ribosomal protein L18a (RPL18a), Ca2+/Calmodulin-dependent protein kinase I (CaMKI), creatine transporter (CRTTR), CDM protein (CDM), adrenoleukodystrophy protein >
7058	19749	328612	2.28	9.7E-01	U28716.1	NT	Drosophila melanogaster sodium channel protein (para) gene, exons 9,10,11,12 and optional segments b, c, d and e, partial cds
8401	21034	34230	1.68	9.7E-01	AF149112.1	NT	Triticum aestivum stripe rust resistance protein Yr10 (Yr10) gene, complete cds
8407	21100	34236	1.3	9.7E-01	M80544.1	NT	Salmonella typhimurium adenine-methyltransferase (mod) and restriction endonuclease (res)
11123	23792		3.94	9.7E-01	BF511298.1	EST_HUMAN	UH+BI-8e1-e-07-0-U1.1 NC1_C3AP_Su88 Homo sapiens cDNA clone IMAGE:3085110 3'
4225	17161	29781	1.5	9.8E-01	AW788874.1	EST_HUMAN	PW2-UM0033-240300-005-12 UM0033 Homo sapiens cDNA
5687	18462	31376	3.77	9.8E-01	Z70558.1	NT	Panvirus B19 DNA, patient C, genome position 2448-2894
5687	18462	31377	3.77	9.8E-01	Z70558.1	NT	Panvirus B19 DNA, patient C, genome position 2448-2894
6848	19410	32224	0.61	9.6E-01	297341.2	NT	Arabidopsis thaliana DNA chromosome 4, ESSA T-FC4 contig fragment No. 6
2291	20895		2.33	9.6E-01	X65275.1	NT	P.falciparum complete gene map of Plasmodium DNA (IR-A)
8750	21442	34689	0.59	9.6E-01	L81138.1	NT	Rettis nonverbal (strain R21) Rsp21 gene, complete cds
11503	24104	37416	3.47	9.6E-01	AV752605.1	EST_HUMAN	AV752605 NP0 Homo sapiens cDNA clone NPDBAG03 5'
11503	24104	37417	3.47	9.6E-01	AV752605.1	EST_HUMAN	AV752605 NP0 Homo sapiens cDNA clone NPDBAG03 5'
11552	24655		1.92	9.6E-01	11421722 NT	NT	Homo sapiens centrosomal protein 2 (CEP2), mRNA
12558	25301	30710	2.18	9.6E-01	U91423.1	NT	Sphyrna tiburo NADH dehydrogenase subunit 2 (NADH2) gene, mitochondrial gene encoding mitochondrial protein, partial cds
2480	16198	27838	1.05	9.6E-01	7705581 NT	NT	Homo sapiens CG1-125 protein (LOC51003), mRNA
2573	15382	28122	0.97	9.5E-01	Q02834	SWISSPROT	ENDOGLICANASE I PRECURSOR (EGI) (ENDO-1,4-BETA-GLUCANASE) (CELLULASE I)
3762	16514	28150	2.04	9.5E-01	BE502340.1	EST_HUMAN	601675638F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3858473 5'
3762	16514	28151	2.04	9.5E-01	BE502340.1	EST_HUMAN	601675638F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3858473 5'
8899	21690	34730	0.69	9.5E-01	AI180162.1	EST_HUMAN	cd57d07_x1 Score= tests_NHT Homo sapiens cDNA clone IMAGE:1733881 3'
9003	21693	34843	1.05	9.5E-01	AW8851102.1	EST_HUMAN	RC1-CT0285-241199-011-b02 CT0285 Homo sapiens cDNA

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Table 4  
Single Exon Probes Expressed In Brain

Probe SEQ ID NO:	Exn SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
11209	23872	37159	1.68	9.5E-01	BF216771.1	EST_HUMAN	601885163 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:4103630 5'
11429	23186	38427	2.42	9.5E-01	AW282788.1	EST_HUMAN	U1-H-BI2_esp4-03-0-UJ.st NCI_CGAP_Sub4 Homo sapiens cDNA clone IMAGE:2727677 3'
11785	24385	37718	1.55	9.5E-01	T87204.1	EST_HUMAN	ya53dd04.e1 Severe fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:66631 3'
3196	15959		3.33	9.4E-01	AF065890.1	NT	Bartonella claritidgeae RNA polymerase beta subunit (rpob) gene, partial cds
3212	15975		2.08	9.4E-01	AF080595.1	NT	Pimpinella brachycarpa zinc finger protein (ZFP1) mRNA, complete cds
8764	21456	34698	0.87	9.4E-01	M90724.1	NT	Human Fc-gamma-receptorIIA (FCGR2A) gene, exon 4
12202	24670		1.82	9.4E-01	BE781251.1	EST_HUMAN	601468703 NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3868928 5'
12557	25219		1.79	9.4E-01	11418857	NT	Homo sapiens epidermal growth factor receptor (avan erythroleastic leukemia viral (v-erb-b) oncogene homolog) (EGFR), mRNA
1723	14468		1.05	9.3E-01	AF242382.1	NT	Homo sapiens phytanoyl-CoA hydroxylase (PHYH) gene, exon 5
2840	15331	28095	1.38	9.3E-01	BE5071172.1	EST_HUMAN	RC8-BT0503-271158-001-801 BT0503 Homo sapiens cDNA
4015	16761	28388	0.88	9.3E-01	M20219.1	NT	Bovine papillomavirus type 2, complete genome
4016	16761	28389	0.88	9.3E-01	M20219.1	NT	Bovine papillomavirus type 2, complete genome
5505	18303	31204	1.56	9.3E-01	AF213984.1	NT	Homo sapiens nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (NFkB1) gene, complete cds
5592	18388	31288	3.89	9.3E-01	L36189.1	NT	Spodoptera frugiperda methionine tetrahydrofolate dehydrogenase mRNA, complete cds
7806	20681	33785	1.85	9.3E-01	AA847040.1	EST_HUMAN	ce098d03.e1 NCI_CGAP_Ov2 Homo sapiens cDNA clone IMAGE:1385337
8713	21405		1.04	9.3E-01	AF061981.1	NT	Xenopus laevis CCCH zinc finger protein C3H-2 (C3H-2) mRNA, complete cds
8835	21527	34673	0.95	9.3E-01	AL161534.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 34
12281	24970		3.12	9.3E-01	AF271207.1	NT	Aedes triseriatus putative large subunit ribosomal protein rp-24 mRNA, complete cds
12802	25049					NT	Homo sapiens chromosome Xq28 melanoma antigen family A2a (MAGEA2a), melanoma antigen family A12 (MAGEA12), melanoma antigen family A2b (MAGEA2b), melanoma antigen family A3 (MAGEA3), cathepsin (CALT), NAD(P)H dehydrogenase-like protein (NSDHL), and L1>
3223	15695	28848	2.83	9.2E-01	BE622702.1	EST_HUMAN	6014413871 NIH_MGC_72 Homo sapiens cDNA clone IMAGE:39161784.3'
4822	17553		0.97	9.2E-01	BF128873.1	EST_HUMAN	6018178147 NIH_MGC_88 Homo sapiens cDNA clone IMAGE:4041363 5'
5531	18428		1.15	9.2E-01	7108410	NT	Mus musculus solute carrier family 30 (zinc transporter), member 4 (SLC30A4), mRNA
5898	18683	31681	7.38	9.2E-01	BF037883.1	EST_HUMAN	601461153 NIH_MGC_86 Homo sapiens cDNA clone IMAGE:3884861 5'
6537	18302	32308	0.61	9.2E-01	MB4703.1	NT	N.crasca valy-tRNA synthetase (cyt-20Un-3) gene
3660	22213	36399	0.92	9.2E-01	AL161565.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 85
38448	22200	38496	1.07	9.2E-01	6871677	NT	Mus musculus carbonic anhydrase 4 (Car4), mRNA
10165	22813	36031	3.16	9.2E-01	11430983	NT	Homo sapiens lysosomal arylsulphatase-like protein 1 (LALP1), mRNA
10315	22682	36173	1.9	9.2E-01	BF688251.1	EST_HUMAN	7056906.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:3578218 3' similar to SWNU5M_TRYBB
10543	23239	38473	1.63	9.2E-01	BE568811.1	EST_HUMAN	601334945 NIH_MGC_39 Homo sapiens cDNA clone IMAGE:3688714 5'

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Table 4  
Single Exon Probes Expressed In Brain

Probe SEQ ID NO:	Exam SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
11722	24316	37639	1.79	9.2E-01	BF32402.1	EST_HUMAN	60182032F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4052018 5'
1621	14368	27067	1.88	9.1E-01	T88675.1	EST_HUMAN	ye52f01_s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:121369 3' similar to contains Alu repetitive element
2120	14851		2.78	9.1E-01	8922056 NT	Homo sapiens hypothetical protein FLJ20048 (FLJ20048), mRNA	
3220	15883	28814	1.15	9.1E-01	T28418.1	EST_HUMAN	AB200G8R Infant brain, LLNL array of Dr. M. Soares 1NIB Homo sapiens cDNA clone LLAB200G8 5'
3220	15903	28815	1.15	9.1E-01	T28418.1	EST_HUMAN	AB200G8R Infant brain, LLNL array of Dr. M. Soares 1NIB Homo sapiens cDNA clone LLAB200G8 5'
6075	18854	31821	1.28	9.1E-01	L36033.1	NT	Human pre-B cell stimulating factor homologue (SDF1b) mRNA, complete cds
8413	18181	32180	3.53	9.1E-01	Q81704	SWISSPROT	INTER-ALPHA-TRYPSIN INHIBITOR HEAVY CHAIN H3 PRECURSOR (III HEAVY CHAIN H3)
7475	20148	33241	17.82	9.1E-01	AA806623.1	EST_HUMAN	cb71g08.s1 NCI CGAP_GCB1 Homo sapiens cDNA clone IMAGE:1336882 3'
7837	20322	33410	2.34	9.1E-01	U72895.1	NT	Rattus norvegicus Rab3 GDP/GTP exchange protein mRNA, complete cds
10075	22723	36840	0.45	9.1E-01	P38432	SWISSPROT	P80-COILIN
122281	25284		27.98	9.1E-01	AF050113.1	NT	Home sapiens uncoupling protein-3 (UCP3) gene, complete cds
43446	17085	28714	2.08	9.0E-01	AF088810.1	NT	Home sapiens neutrophil III-alpha gene, partial cds
7281	18974	33052	0.72	9.0E-01	U42547.1	NT	Danio rerio LM class homeodomain protein (lim5) mRNA, complete cds
7321	20004		1.18	9.0E-01	D38821.1	NT	Xenopus laevis genes for alkaline phosphatase, complete cds
9249	21928	35100	0.49	9.0E-01	AF088781.1	NT	Danio rerio semaphorin Z1a mRNA, complete cds
5510	18408	31318	2.68	8.9E-01	AF026198.1	NT	Fugu rubripes neural cell adhesion molecule L1 homolog (L1-CAM) gene, complete cds; putative protein 1 (PUT1) gene, partial cds; mitosis-specific chromosome segregation protein SMC1 homolog (SMC1) gene, complete cds; and calcium channel alpha-1 subunit
6154	18831		1.38	8.9E-01	X606986.1	NT	Rabbit MHC fragment RLA-DF DNA
8925	21018	34154	0.71	8.9E-01	AF259887.1	NT	Oithona nana cytochrome-c oxidase subunit I (cox1) gene, partial cds; mitochondrial gene for mitochondrial product
11787	24377	37707	2.51	8.9E-01	AE003944.1	NT	Xylella fastidiosa, section 90 of 229 of the complete genome
12138	24627		2.86	8.9E-01	AE002186.2	NT	Chlamydophila pneumoniae AR39, section 21 of 94 of the complete genome
12762	25343		2.51	8.9E-01	A1150836.1	EST_HUMAN	qb84d08.x1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:1704879 3'
4505	17240	28873	3.82	8.8E-01	O28350	SWISSPROT	PUTATIVE F420-DEPENDENT NADP REDUCTASE
6289	16094	30765	0.67	8.8E-01	AF310617.1	NT	Pseudorabies virus Ea glycoprotein M gene, complete cds
10131	22779	36892	0.83	8.8E-01	7656978	NT	Homo sapiens cell death-inducing DFFA-like effector B (CIDEB), mRNA
11018	23890	36953	4.98	8.8E-01	Z28337.1	NT	M aeruginosa (HUB 5-2-4) DNA from plasmid pMA1
11968	26382		1.8	8.8E-01	D80911.1	NT	Synochocystis sp. PCC6803 complete genome, 13/27, 1576583-1719843
452	13238	25877	1.54	8.7E-01	AF089332	NT	Homo sapiens SOS1 (SOS1) gene, partial cds
2401	15122	27859	1.07	8.7E-01	6901883	NT	Homo sapiens AT-binding transcription factor 1 (ATBFI), mRNA

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**Single Exon Probes Expressed in Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
2877	15644	28288	6.05	8.7E-01	AA595383.1	EST_HUMAN	nn05311.s1 NCI_CGAP_Pt4.1 Homo sapiens cDNA clone IMAGE:1076877
4948	17873			3.17	8.7E-01 AF121970.1	NT	Pseudomonas aeruginosa topoisomerase (top), putative transcriptional regulatory protein OhbR (ohbR), ortho-halobenzene 1,2-dioxygenase beta-ISPF protein OhbA (ohbA), OhbC (ohbC), OhbD (ohbD), and putP dioxygenase alpha-ISPF protein OhbB (ohbB), and putP
5102	17820			0.97	8.7E-01 AJ28805.1	NT	Homo sapiens partial Lgals3 gene for galectin-8, exon 3
7839	20634	33761	0.62	8.7E-01 AW897335.1	EST_HUMAN	RC4-NIN0057-120500-013-007 NIH0057 Homo sapiens cDNA	
8829	21520	34665	0.69	8.7E-01 AI239458.1	EST_HUMAN	qhs6e06.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1846788 3'	
8828	21520	34666	0.69	8.7E-01 AI239456.1	EST_HUMAN	qhs6e06.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1846788 3'	
9838	22290	35453	1.57	8.7E-01 AE004933.1	NT	Pseudomonas aeruginosa PA01, section 524 of 526 of the complete genome	
10202	22850	36065	0.61	8.7E-01 BF570168.1	EST_HUMAN	602185541T1 NIH_MGC_45 Homo sapiens cDNA clone IMAGE:4309808 3'	
10202	22850	36066	0.61	8.7E-01 BF570169.1	EST_HUMAN	602185541T1 NIH_MGC_45 Homo sapiens cDNA clone IMAGE:4309808 3'	
10735	23422	36665	5.25	8.7E-01 BF563070.1	EST_HUMAN	QV0-NN1021-100800-337-003 NN1021 Homo sapiens cDNA	
11739	24332	37657	5.47	8.7E-01 BF107364.1	EST_HUMAN	601823684R1 NIH_MGC_79 Homo sapiens cDNA clone IMAGE:4043564 3'	
11739	24332	37658	5.47	8.7E-01 BF107364.1	EST_HUMAN	601823684R1 NIH_MGC_79 Homo sapiens cDNA clone IMAGE:4043564 3'	
482	13247		1.75	8.6E-01 X17012.1	NT	Ref IGFII  gene for insulin-like growth factor II	
838	13608	26279	3.45	8.6E-01 W69089.1	EST_HUMAN	zH44sd3.r1 Soares_fetal heart_Nib-H19W Homo sapiens cDNA clone IMAGE:343516 5'	
2298	14994	27733	0.93	8.6E-01	4603210 NT	Homo sapiens cytochrome P450, subfamily XXVIIA (steroid 27-hydroxylase, cerebratilinicus xanthomatosis), polypeptide 1 (CYP27A1b) mRNA	
36098	16361	29003	0.85	8.6E-01 AL161865.2	NT	Aribidopsis thaliana DNA chromosome 4, contig fragment No. 65	
3792	16534	29172	1.55	8.6E-01 U49724.1	NT	Drosophila melanogaster mrfn (Mrfn) mRNA, complete cds	
58098	18597	31524	10.86	8.6E-01 X80547.1	NT	Chicken lipoprotein lipase gene	
59018	18597	31525	10.88	8.6E-01 X80547.1	NT	Chicken lipoprotein lipase gene	
65009	18372	32285	2.08	8.6E-01 AF143732.1	NT	Grus canadensis recombination activating protein 1 (RAG-1) gene, partial cds	
65009	18372	32286	2.08	8.6E-01 AF143732.1	NT	Grus canadensis recombination activating protein 1 (RAG-1) gene, partial cds	
7427	20104		0.78	8.6E-01 AE000897.1	NT	Halobacter pylori 26895 section 88 of 134 of the complete genome	
7828	20523		1.12	8.6E-01 AP001518.1	NT	Bacillus halodurans genomic DNA, section 12/14	
7841	20836	33763	0.55	8.6E-01 AF077837.1	NT	Drosophila melanogaster collapsin response mediator protein (CRMP) mRNA, complete cds	
8585	22238		0.40	8.6E-01 AE000878.1	NT	Araeaogobius fujigae section 128 of 172 of the complete genome	
126518	25144		1.35	8.6E-01 AL112162.1	NT	Batrachis ciliaris strain T4 cDNA library under conditions of nitrogen deprivation	
66226	18388	32401	0.95	8.5E-01 AF1685214.1	NT	Bacteriophage D3, complete genome	
7425	20102	33189	2.51	8.5E-01 BE542612.1	EST_HUMAN	60108710771 NIH_MGC_10 Homo sapiens cDNA clone IMAGE:3453605 5'	
8317	21010	34147	0.78	8.5E-01 P08601	SWISSPROT	SEGMENTATION PROTEIN PAIRED	
8317	21010	34148	0.78	8.5E-01 P08601	SWISSPROT	SEGMENTATION PROTEIN PAIRED	
8402	21056	34231	0.67	8.5E-01 AJ24S213.1	NT	Homo sapiens partial 5-HT4 receptor gene, exons 2 to 5	

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar BLAST E Value	Top Hit No.	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
10249	22896	36105	1.17	8.5E-01	AB006799.1	NT	Cyanidium caldarium gene for SigC, complete cds	
10249	22896	36106	1.17	8.5E-01	AB006799.1	NT	Cyanidium caldarium gene for SigC, complete cds	
12278	25286		2.24	8.5E-01	11418543	NT	Homo sapiens human immunodeficiency virus type 1 enhancer-binding protein 1 (HIVEP1), mRNA	
4702	17438	30067	0.73	8.4E-01	AF083976.2	NT	Fowl adenovirus 8, complete genome	
6406	25068	30910	2.29	8.4E-01	L78728.1	NT	Human fibroblast growth factor receptor 3 (FGFR3) gene, intron 7	
6406	25068	30911	2.28	8.4E-01	L78728.1	NT	Human fibroblast growth factor receptor 3 (FGFR3) gene, intron 7	
7708	20372	33485	0.63	8.4E-01	AF051142.1	NT	Manestra brassicae pheromone binding protein 2 precursor (FBP2) mRNA, complete cds	
8898	22508		2.68	8.4E-01	AJ248287.1	NT	Pyrrococcus abyssi complete genome, segment 5/6	
724	13498	26151	2.8	8.3E-01	M63437.1	NT	Thermus thermophilus cytochrome c-552 (cytC) and CycB (cycB) genes, complete cds	
30891	15898	28497	2.90	8.3E-01	AL161606.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 18	
31780	18542	28177	0.79	8.3E-01	AB010879.1	NT	Nicotiana tabacum mRNA for chloroplast ribosomal protein L10, complete cds	
36933	16741	28375	3.35	8.3E-01	Y19177.1	NT	Streptomyces antibioticus polyketide biosynthetic gene cluster	
5187	17985	30511	2.41	8.3E-01	AL161640.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 40	
9568	22221		4.51	8.3E-01	AJ791852.1	EST_HUMAN	m01f12-5 NCI_CGAP_Co9 Homo sapiens cDNA clone IMAGE-1076495 5' similar to contains ThR-H1 THR repetitive element	
10010	22658	35872	1.27	8.3E-01	AF098070.1	NT	Drosophila melanogaster Usl1 homolog mRNA, complete cds	
10118	22768	35873	3.46	8.3E-01	AF108133.1	NT	Mus musculus neuro-d4 gene, exons 3 through 12 and partial cds	
10572	23267	36506	3.35	8.3E-01	AE000903.1	NT	Methanobacterium thermoautotrophicum from bases 1270510 to 1283409 (section 109 of 148) of the complete genome	
10590	23284		2.03	8.3E-01	7212472	NT	Phytophthora infestans mitochondrial, complete genome	
11274	23935	37227	2	8.3E-01	AF020503.1	NT	Homo sapiens FRAS2 common fragile region, diastenosine triphosphate hydrolase (FHIT) gene, exon 5	
20445	14778	27506	2.3	8.2E-01	AB000489.1	NT	Rattus norvegicus mRNA for RPHO-1, complete cds	
2083	14816		1.31	8.2E-01	AF146589.1	NT	Mus musculus trophinin (Tnn) gene, complete cds	
2888	15395		1.08	8.2E-01	AW376860.1	EST_HUMAN	IL-3-C10219-181189-031-008 CT0219 Homo sapiens cDNA	
6876	16583	32631	0.75	8.2E-01	AJ010142.1	NT	Amanita muscaria mRNA for SCII125 protein	
6797	18541	32569	3.49	8.2E-01	AW378433.1	EST_HUMAN	CM4-HT0243-081189-037-001 HT0243 Homo sapiens cDNA	
7169	25108		3.2926	4.74	8.2E-01	Z12128.1	S.cerevisiae MET, LEU4, and POL1 genes encoding MET4 protein, alpha-sopropylmalate (alpha-[PM]) synthetase (partial), and DNA polymerase alpha (partial)	
8343	21036	34173	0.59	8.2E-01	BE283145.1	EST_HUMAN	60114488572 NIH_MGC_18 Homo sapiens cDNA clone IMAGE-3160412 5'	
8924	22572	35770	0.65	8.2E-01	AB014530.1	NT	Homo sapiens mRNA for KIAA0630 protein, partial cds	
8859	22607	35812	1.37	8.2E-01	AF052859.1	NT	Homo sapiens thioredoxin-related protein mRNA, complete cds	
10123	22771	35985	0.59	8.2E-01	AE223888.1	NT	Oncorhynchus tshawytscha Isolate T-20 somatotrophin precursor gene, exon 1	
10123	22771	35988	0.59	8.2E-01	AF223888.1	NT	Oncorhynchus tshawytscha Isolate T-20 somatotrophin precursor gene, exon 1	

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Meet Similar (T <sub>ce</sub> ) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
10283	22831	36145	3.65	8.2E-01	Q8JL70	SWISSPROT	MCKUSICK-KAUFMAN/BARDET-BIEDL SYNDROMES PUTATIVE CHAPERONIN
10283	22831	36148	3.65	8.2E-01	Q8JL70	SWISSPROT	MCKUSICK-KAUFMAN/BARDET-BIEDL SYNDROMES PUTATIVE CHAPERONIN
11841	24238	37562	4.65	8.2E-01	L10127.1	NT	Molluscum contagiosum virus type 1 ORF1 and ORF2 DNA
11735	24828	37652	6.38	8.2E-01	P10383	SWISSPROT	OVARIAN TUMOR LOCUS PROTEIN
11740	24333	37659	6.02	8.2E-01	H87398.1	EST_HUMAN	yw1402.1 Scores_placenta_86weeks_2N18P58eW Homo sapiens cDNA clone IMAGE:2521955
12298	24723	31054	2.37	8.2E-01	AJ001261.1	NT	similar to gb:M36072 60S RIBOSOMAL PROTEIN L7A (HUMAN);
2762	15467		1.08	8.1E-01	AF181839.1	NT	Mus musculus mRNA for NIPSNAP2 protein
3451	16207	28857	3.08	8.1E-01	AF055088.1	NT	Mus musculus TANK binding kinase TBK1 (TBK1) mRNA, complete cds
3451	16207	28858	3.08	8.1E-01	AF055088.1	NT	Homo sapiens NiHC class 1 region
4883	17592		0.74	8.1E-01	AF202634.1	NT	Drosophila melanogaster Na/K-ATPase beta subunit isoform 4 (JYbeta2) mRNA, complete cds
62223	18997	31973	0.84	8.1E-01	U18790.1	NT	Mus musculus putative collagen alpha-2(XI) chain (COL11A2) gene, partial cds
6526	19292	32285	2.68	8.1E-01	Q13491	SWISSPROT	NEURONAL MEMBRANE GLYCOPROTEIN M6-B
6526	19292	32286	2.68	8.1E-01	Q13491	SWISSPROT	NEURONAL MEMBRANE GLYCOPROTEIN M6-B
7226	19914	32987	0.78	8.1E-01	AB007877.1	NT	Homo sapiens KIAA0417 mRNA, complete cds
7412	20089	33173	0.68	8.1E-01	O47477	SWISSPROT	CYTOKROME B
7811	20506	33638	0.75	8.1E-01	AF022713.2	NT	Drosophila melanogaster putative inorganic phosphate cotransporter (Pico) gene, partial cds; putative sodium channel (NaCh) and putative amylase-related protein (Amyre) genes, complete cds; and putative serine-enriched protein (gprs) gene, partial cds
7811	20506	33639	0.75	8.1E-01	AF022713.2	NT	Drosophila melanogaster putative inorganic phosphate cotransporter (Pico) gene, partial cds; putative sodium channel (NaCh) and putative amylase-related protein (Amyre) genes, complete cds; and putative serine-enriched protein (gprs) gene, partial cds
8507	21189	34344	0.58	8.1E-01	AF001517.1	NT	Bacillus halodurans genomic DNA, section 11/14
8507	21189	34345	0.58	8.1E-01	AF001517.1	NT	Bacillus halodurans genomic DNA, section 11/14
86688	21330	34507	1.08	8.1E-01	AW242847.1	EST_HUMAN	xn01h03_x1_NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2682469 3' similar to SWLYAR_MOUSE_Q08288 CELL GROWTH REGULATING NUCLEOLAR PROTEIN; contains MER22_b1 PTR5_repetitive element;
10025	22673	35888	0.7	8.1E-01	P08425	SWISSPROT	PROBABLE E4 PROTEIN
10311	22858	36174	0.6	8.1E-01	N84541.1	EST_HUMAN	KK9872 Human fetal heart, Lambda ZAP Express Homo sapiens cDNA clone KK9872 5' similar to ESTCLONE C-0PE11)
11484	24067	37374	2.63	8.1E-01	BE838558.1	EST_HUMAN	RC0-TN0080-220800-025-010 TN0080 Homo sapiens cDNA
11484	24067	37375	2.63	8.1E-01	BE838558.1	EST_HUMAN	RC0-TN0080-220800-025-010 TN0080 Homo sapiens cDNA
12022	24550	31110	1.57	8.1E-01	AE001711.1	NT	Thermatoga maritima section 23 of the complete genome
172	12385		3.49	8.0E-01	AJ271510.1	NT	Staphylococcus aureus partial pta gene for phosphatid acyltransferase allele 15

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
282	13089	25720	13.81	8.0E-01	AJ132772.1	NT	<i>Bos taurus fib and rtf genes</i>
1595	14341	27031	1.12	8.0E-01	8394087	NT	<i>Rattus norvegicus protease (procathepsin, macrophain) 28 subunit, alpha (Psmce1), mRNA</i>
2029	14764		1.91	8.0E-01	BF530862.1	EST_HUMAN	602072473F1 NCI_CGAP_Bm87 Homo sapiens cDNA clone IMAGE:4215091 5'
3076	15844	28484	1.2	8.0E-01	AF427897.1	NT	<i>Semini bivalvulae olfactory receptor (SBOZ7) gene, partial cds</i>
3307	16067	28716	1.35	8.0E-01	AB006198.1	NT	<i>Mus musculus gene for avian/duchii chaperonin, complete cds</i>
3680	18443		1.52	8.0E-01	AL1622758.2	NT	<i>Neisseria meningitidis serogroup A strain 22481 complete genome; segment 77</i>
4498	17232	29862	8.05	8.0E-01	X83739.2	NT	<i>G. gallus mRNA for nicotinic acetylcholine receptor (MACHR) beta 3 subunit</i>
7889	20584		2.25	8.0E-01	AW901489.1	EST_HUMAN	RC0-NIN1012-270300-021-H05 NN1012 Homo sapiens cDNA
8423	21110	34254	0.98	8.0E-01	Y11085.1	NT	<i>Rice stripe virus RNA 3</i>
10876	235563	36803	2.78	8.0E-01	Q9Z783	SWISSPROT	<i>CREB-BINDING PROTEIN</i>
4411	13227	25870	1.16	7.9E-01	D11478.1	NT	<i>Lymphotoxin delta nuclear polyhedrosis virus gene for DNA polymerase, complete cds</i>
698	13473		1.14	7.9E-01	AE002180.1	NT	<i>Ureaplasma urealyticum section 31 of 59 of the complete genome</i>
1600	14343		22.89	7.9E-01	AB040885.1	NT	<i>Homo sapiens mRNA for KIAA1482 protein, partial cds</i>
1652	14393		1.2	7.9E-01	UR2739.1	NT	<i>Haemophilus influenzae Rd section 54 of 163 of the complete genome</i>
2269	14989	27726	5.86	7.9E-01	AB004816.1	NT	<i>Oryctolagus cuniculus mRNA for mitsugumin28, complete cds</i>
2260	14987	27727	2.4	7.9E-01	AF130489.1	NT	<i>Danio rerio Tp4-associated protein Tap1A (Tap1A) mRNA, complete cds</i>
3568	162622	28916	3.01	7.9E-01	AF228884.1	NT	<i>Gallus gallus SOX8 transcription factor (SOX8) mRNA, complete cds</i>
4288	17008		0.85	7.9E-01	BE283612.1	EST_HUMAN	601912033F1 NIH_MCGC_7 Homo sapiens cDNA clone IMAGE:3535785 5'
4572	17307	28635	1.13	7.9E-01	6753746	NT	<i>Mus musculus embigin (Emb), mRNA</i>
4672	17307	28836	1.13	7.9E-01	6753745	NT	<i>Mus musculus embigin (Emb), mRNA</i>
6252	19029		0.57	7.9E-01	D28145.1	NT	<i>Human mRNA for prostacyclin synthase, complete cds</i>
8508	20703	33831	2.79	7.9E-01	X90986.1	NT	<i>P. sativum GR gene</i>
9447	22124	36804	4.04	7.9E-01	U01812.1	NT	<i>Glandula lambla variant-specific surface protein G3M-B (vsgG3M-B) mRNA, partial cds</i>
9949	22597	35801	4.47	7.9E-01	F19719	SWISSPROT	<i>SMALL HYDROPHOBIC PROTEIN</i>
9891	22639	35849	0.51	7.9E-01	AV700860.1	EST_HUMAN	AV700860 GKC Homo sapiens cDNA clone GKCDRE123'
10408	23054	36271	1.24	7.9E-01	AB000531.1	NT	<i>Streptococcus mutans DNA for sigma 42 protein, dTDP-4-keto-L-rhamnose reductase, complete cds</i>
10516	23162	36369	0.52	7.9E-01	P16305	SWISSPROT	<i>DYNEIN HEAVY CHAIN (DYHC)</i>
10629	23609		2.74	7.9E-01	7662471	NT	<i>Homo sapiens KIAA1072 protein (KIAA1072), mRNA</i>
11173	23840	37123	2.02	7.9E-01	P18022	SWISSPROT	<i>NEURAL-CADHERIN PRECURSOR (N-CADHERIN)</i>
858	13625		2.24	7.8E-01	Z43785.1	EST_HUMAN	HSC1KH041 normalized infant brain cDNA Homo sapiens cDNA clone c-1Kh04
2273	14999	27737	7.47	7.8E-01	AW888667.1	EST_HUMAN	EST371637 MAGE resequences, MAGF Homo sapiens cDNA
4653	17387	30020	0.73	7.8E-01	U87305.1	NT	<i>Rattus norvegicus transmembrane receptor Utrophin 1 mRNA, complete cds</i>
6578	18760	31724	2.28	7.8E-01	AF115856.1	NT	<i>Sphingon punctatus alpha endopeptidase mRNA, partial cds</i>

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
6124	18902	31871	0.88	7.8E-01	P05231	SWISSPROT	INTERLEUKIN-6 PRECURSOR (IL-6) (B-CELL STIMULATORY FACTOR 2) (BSF-2) (INTERFERRON BETA-2) (HYBRIDOMA GROWTH FACTOR)
6371	18140	32136	0.63	7.8E-01	AL445068.1	NT	Theroplasma acidophilum complete genome; segment 4/5
8389	21082	34216	1.02	7.8E-01	BF108927.1	EST_HUMAN	Discoidium fusca cDNA clone IMAGE:3525178 3'
9133	21821	34887	1.3	7.8E-01	Y10168.1	NT	Discoidium fusca cDNA clone IMAGE:3525178 3'
9231	21910	36083	0.51	7.8E-01	48268873	NT	Homo sapiens nucleophillin 21 kDa (CAIN) (NUP214), mRNA
10024	22672		0.97	7.8E-01	Q25462	SWISSPROT	MUSCLE CALCIUM CHANNEL ALPHA-1 SUBUNIT (MDL-ALPHA1)
12271	25273		2.5	7.8E-01	L29280.1	NT	Arabidopsis thaliana 1-amino-1-cyclopentanecarboxylate synthase (ACS5) gene, complete cds
139	12854	25598	7.61	7.7E-01	AF184345.1	NT	Lycorestanin ADP-glucosidase large subunit (AGP-L1) mRNA, complete cds
709	13483		2.26	7.7E-01	AF050157.1	NT	Mus musculus major histocompatibility locus class II region: major histocompatibility protein class II alpha butyrophilin-like (NG9), butyrophilin-1p
2717	15424	28163	2.21	7.7E-01	O33915	SWISSPROT	CITRATE SYNTHASE
3351	16111		0.84	7.7E-01	8388408	NT	Homo sapiens UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-acetylgalactosaminyltransferase 7 (GalNAc-T7) (GALNACT7), mRNA
3686	16340	28985	3.98	7.7E-01	AF118055.1	NT	Homo sapiens PRO19875 mRNA, complete cds
4385	17103	29738	3.98	7.7E-01	AF189488.1	NT	Colomix columnica sub-species japonica beta-actin mRNA, partial cds
4386	17103	29739	3.98	7.7E-01	AF189488.1	NT	Colomix columnica sub-species japonica beta-actin mRNA, partial cds
5473	18272	31165	1.45	7.7E-01	P18653	SWISSPROT	RAFFINOSE INVERTASE (INVERTASE)
6473	18272	31168	1.45	7.7E-01	P18653	SWISSPROT	RAFFINOSE INVERTASE (INVERTASE)
6868	18853	31594	0.85	7.7E-01	R08800.1	EST_HUMAN	Myo 402.51 Soares fetal liver spleen (NF1S) Homo sapiens cDNA clone IMAGE:127755 3'
9744	22395	35890	0.51	7.7E-01	AB021134.1	NT	Daphnia magna hemoglobin gene cluster (dhb3, dhb1 and dhb2 genes), complete cds
12161	24844		4.55	7.7E-01	11497621	NT	Archaeoglobus fulgidus, complete genome
6008	18789	31751	4.88	7.8E-01	AF058510.1	NT	Arabidopsis thaliana 3-methylcrotonyl-CoA carboxylase non-biotinylated subunit (MCCB) mRNA, complete cds
6008	18789	31752	4.88	7.8E-01	AF058510.1	NT	Arabidopsis thaliana 3-methylcrotonyl-CoA carboxylase non-biotinylated subunit (MCCB) mRNA, complete cds
6425	19193	32189	0.81	7.8E-01	P27938	SWISSPROT	MATING-TYPE PROTEIN A:ALPHA 74
6761	17920	30556	0.94	7.8E-01	AI258398.1	EST_HUMAN	eq14b12.51 Stanley Frontal NS pool 2 Homo sapiens cDNA clone IMAGE:2033879
6761	17920	30585	0.94	7.8E-01	AI258399.1	EST_HUMAN	eq14b12.51 Stanley Frontal NS pool 2 Homo sapiens cDNA clone IMAGE:2033879
6951	19433	32449	0.88	7.8E-01	U72487.1	NT	Reptilia norvegicus calcium-independent alpha-leetroxin receptor mRNA, complete cds

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar BLAST E Value	(Top) Hit No.	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
7984	20859	33784	1.38	7.6E-01	AF146793.2	NT	Mus musculus neuregulin U precursor (Nmju) gene, partial cds; BPhLP (Tphlp) gene, partial cds; CLOCK (Clock) gene, complete cds; PFT27 (Pf27) gene, complete cds; and HSAR (Hsar) gene, complete cds	
8026	20721	33852	1.88	7.6E-01	6857752	NT	Mus musculus adillin (Adm1-pending), mRNA	
8026	20721	33853	1.88	7.6E-01	6857752	NT	Mus musculus adillin (Adm1-pending), mRNA	
8868	21557	34703	0.74	7.6E-01	6763577	NT	Mus musculus cytochrome P450, 2B9, phenobarbital inducible, type a (Cyp2b9), mRNA	
9170	21849	35015	6.03	7.6E-01	P30372	SWISSPROT	MUSCARINIC ACETYLCHOLINE RECEPTOR M2	
9179	21849	35016	5.03	7.6E-01	P30372	SWISSPROT	MUSCARINIC ACETYLCHOLINE RECEPTOR M2	
11330	24021	37325	2.68	7.6E-01	X86347.1	NT	H aspera mRNA for neurofilament NF70	
11330	24021	37326	2.68	7.6E-01	X86347.1	NT	H aspera mRNA for neurofilament NF70	
11711	24306		3.64	7.6E-01	AL161592.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 88	
11831	24489		3.73	7.6E-01	AB020702.1	NT	Homo sapiens mRNA for KIAA0895 protein, partial cds	
500	13284		1.44	7.6E-01	AL163301.2	NT	Homo sapiens chromosome 21 segment HS21C101	
570	13351	25979	1.23	7.5E-01	AF020593.1	NT	Homo sapiens FRA3B common fragile region, diadenosine triphosphate hydrolase (FHTT) gene, exon 5	
3354	16114	28769	0.95	7.5E-01	C14203.1	EST_HUMAN	C14203 Clontech human aorta polyA+ mRNA (#8572) Homo sapiens cDNA clone GEN-037E11 5'	
7421	20098	33186	1.01	7.5E-01	AF052730.1	NT	Drosophila melanogaster tyrosine kinase receptor protein (eph) mRNA, complete cds	
11177	23844	37130	1.5	7.5E-01	AB047819.1	NT	Homo sapiens GCMa/GCM1 gene for chorion-specific transcription factor GCMa, complete cds	
12228	24682		4.3	7.5E-01	AF163161.2	NT	Homo sapiens dentin sialophosphoprotein precursor (DSPP) gene, complete cds	
12742	25008	30976	1.46	7.5E-01	AE000823.1	NT	Methanobacterium thermophilicum from bases 317350 to 328782 (section 28 of 148) of the complete genome	
1108	13886	28522	1.78	7.4E-01	AL598146.1	EST_HUMAN	In14b08.x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE-2167577 3' similar to contains Alu repetitive elementcontains element MIR repetitive element :	
2342	15065	27802	0.98	7.4E-01	AB011106.1	NT	Homo sapiens mRNA for KIAA0534 protein, partial cds	
4273	17016	29842	4.73	7.4E-01	AL163246.2	NT	Homo sapiens chromosome 21 segment HS21C048	
7743	20430	33662	1.23	7.4E-01	AL161651.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 51	
7743	20430	33583	1.23	7.4E-01	AL161651.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 51	
8831	21223	34265	0.83	7.4E-01	BF346286.1	EST_HUMAN	602018465F1 NCI CGAP_Bm67 Homo sapiens cDNA clone IMAGE-4164340 5'	
8863	21305		0.76	7.4E-01	U87980.1	NT	Rattus norvegicus leukocyte common antigen receptor (LAR) gene, trans-spliced alternative untranslated exon	
8864	21684	34834	6.95	7.4E-01	BE747603.1	EST_HUMAN	601673026F1 NIH MGIC_9 Homo sapiens cDNA clone IMAGE-3834174 5'	
9054	21743	34901	1.14	7.4E-01	AA187988.1	EST_HUMAN	2P67H101.51 Stratagene endothelial cell S37223 Homo sapiens cDNA clone IMAGE-825297 3' similar to SW:TCPQ_MOUSE_P42932 T-COMPLEX PROTEIN 1, THETA SUBUNIT;	
10302	22349	36164	0.76	7.4E-01	11424833	NT	Homo sapiens NY-REN-45 antigen (LOC51133), mRNA	

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**Table 4**  
**Single Exon Probes Expressed in Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
11685	24260	37682	1.65	7.4E-01	AB021480.2	NT	Oryzias latipes gene for membrane guanylyl cyclase OIGC1, complete cds
11685	24260	37583	1.65	7.4E-01	AB021480.2	NT	Oryzias latipes gene for membrane guanylyl cyclase OIGC1, complete cds
11800	24467		3.62	7.4E-01	6753217	NT	Mus musculus complement component 1 inhibitor (C1inh), mRNA
12008	24542		1.78	7.4E-01	AI472841.1	EST_HUMAN	rat3h01.X1 NCI_CGAP Lym5 Homo sapiens cDNA clone IMAGE:2043985 3'
2889	15765	28413	0.8	7.3E-01	P08710	SWISSPROT	HYPOTHETICAL PROTEIN HKLF1 (IRL1) (TRL1)
4575	17310	28838	0.7	7.3E-01	AE001186.1	NT	Bacilla буддара (section 62 of 70) of the complete genome
4652	17386	30019	4.37	7.3E-01	AF225421.1	NT	Homo sapiens HT017 mRNA, complete cds
5040	17759	30373	1.01	7.3E-01	O49103	SWISSPROT	FERRICIRONOME SIDEWORPHORE PEPTIDE SYNTHETASE
6511	19278	32278	5.92	7.3E-01	I35772.1	NT	Mus musculus antigen (CD72) gene
6511	19278	32277	6.92	7.3E-01	I35772.1	NT	Mus musculus antigen (CD72) gene
6894	25103	32735	0.97	7.3E-01	AJ011418.1	NT	Lycopersicon esculentum mRNA for ubiquitin activating enzyme
7359	20040	33118	0.98	7.3E-01	Z14133.1	NT	D.melanogaster Chc mRNA for clathrin heavy chain
7445	20121	33210	7.94	7.3E-01	M28511.1	NT	V.aliginolyticus sucrose (scrB) gene, complete cds
7445	20121	33211	7.94	7.3E-01	M28511.1	NT	V.aliginolyticus sucrose (scrB) gene, complete cds
11407	24058	37561	3.53	7.3E-01	AA878019.1	EST_HUMAN	222508.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:431798 3'
11407	24058	37562	3.53	7.3E-01	AA878019.1	EST_HUMAN	222508.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:431798 3'
812	13583		3.98	7.2E-01	L28281.1	NT	Rattus norvegicus histone factor-2 kinase (elf-2a) mRNA, complete cds
1950	14885	27398	2.32	7.2E-01	X78140.1	NT	N.tabacum Nelf-4A13 mRNA
2453	16181	27920	1.27	7.2E-01	AB009805.1	NT	Gallus gallus gene for melanocortin 2-receptor, complete cds
3063	15829	28473	1.38	7.2E-01	AF198100.1	NT	Fowl/pox virus, complete genome
3445	16201	28851	2.58	7.2E-01	AF086608.1	NT	Giardia lamblia variant-specific surface protein (vsp417-6) gene, vsp417-6/A-H allele, complete cds
3601	16364	28994	1.08	7.2E-01	AB002307.1	NT	Human mRNA for KIAA0309 gene, partial cds
4040	16785		0.7	7.2E-01	AF108093.1	NT	Homo sapiens IA-2 gene, intron 18
4718	17450	30083	2.65	7.2E-01	D80314.1	NT	messenger ribonucleic acid for sucrose phosphorylase (EC:2.4.1.7)
5075	17794	30410	0.74	7.2E-01	P33066	SWISSPROT	NUCLEOSIDE TRIPHOSPHATASE I (NUCLEOSIDE TRIPHOSPHATE PHOSPHOHYDROLASE I) (NPH I)
7112	19800	32884	0.88	7.2E-01	U69833.1	NT	Solanum tuberosum cold-stress inducible protein (C17) gene, complete cds
8353	21046	34183	1.11	7.2E-01	AF236061.1	NT	Oryctolagus cuniculus RING-finger binding protein mRNA, partial cds
8862	21653		0.48	7.2E-01	AV743773.1	EST_HUMAN	AV743773 CB Homo sapiens cDNA clone CEMAFD08 5'
10239	22887	36100	2.33	7.2E-01	BF670067.1	EST_HUMAN	602118381(F1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:4275381 6'
10639	23330	36568	4.02	7.2E-01	U82623.1	NT	Rattus norvegicus cytochrome b mRNA, complete cds
11104	23774	37049	1.27	7.2E-01	S76838.1	NT	Dbs=Dbs guanine nucleotide exchange factor homolog [mice, 32D murine hemopoietic cell line, mRNA, 3923 nf]

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Table 4

Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (TBLASTN) Hit Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
12422	24798		2.9	7.2E-01 AP000068.1	NT		Aeropynum permix genomic DNA, section 6/7
878	13451	28094	12.73	7.1E-01 D21070.1	NT		Rana catesbeiana mRNA for bullfrog skeletal muscle calcium release channel (ryanodine receptor) alpha isoform (RYR1), complete cds
3059	18826	28470	11.76	7.1E-01 AJ270777.1	NT		Homo sapiens partial TCF-4 gene for T-cell transcription factor-4, exons 15-16
4184	18926	28555	3.18	7.1E-01	7305360 NT		Mitis musculus docephin (Otoc), mRNA
4184	18925	28556	3.18	7.1E-01	7305360 NT		Mitis musculus docephin (Otoc), mRNA
58558	18846	31685	1.63	7.1E-01 BF081034.1	EST_HUMAN	602155498F1 NIH MGC_83 Homo sapiens cDNA clone IMAGE:4296344 5'	
58553	18845	31686	1.63	7.1E-01 BF081034.1	EST_HUMAN	602155498F1 NIH MGC_83 Homo sapiens cDNA clone IMAGE:4296344 5'	
68550	19550	32850	7.68	7.1E-01 U862232.1	NT	Drosophila melanogaster 6-pyruvotetrahydropterin synthase (pt) gene, complete cds	
80891	20785	33818	0.88	7.1E-01 HS4244.1	EST_HUMAN	yp89d09.s1 Scores fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:202861 3'	
86355	21327	34498	0.93	7.1E-01 BE074185.1	EST_HUMAN	RC1-BT0537-301289-011-009 BT0537 Homo sapiens cDNA	
86355	21327	34470	0.93	7.1E-01 BE074185.1	EST_HUMAN	RC1-BT0537-301289-011-009 BT0537 Homo sapiens cDNA	
97555	22408	35693	1.43	7.1E-01 BE04405.1	EST_HUMAN	601498330F1 NIH MGC_70 Homo sapiens cDNA clone IMAGE:3898495 5'	
10309	22956	36172	1.22	7.1E-01 M12861.1	NT	Human T-cell receptor gamma-chain J2 gene	
12211	25205		2.21	7.1E-01 AA421482.1	EST_HUMAN	z06fh1.s1 Scores testis_NHT Homo sapiens cDNA clone IMAGE:731108 3'	
12097	13958	28624	0.98	7.0E-01 AB014514.1	NT	Homo sapiens mRNA for KIAA0014 protein, partial cds	
12097	13958	28625	0.98	7.0E-01 AB014514.1	NT	Homo sapiens mRNA for KIAA0014 protein, partial cds	
2460	15169	27807	1.13	7.0E-01 NB2412.1	EST_HUMAN	y273d07.s1 Scores_multiple_scattered_2NBHMASP Homo sapiens cDNA clone IMAGE:288708 3' similar to contains Alt repetitive element	
2460	15169	27808	1.13	7.0E-01 NB2412.1	EST_HUMAN	y273d07.s1 Scores_multiple_scattered_2NBHMASP Homo sapiens cDNA clone IMAGE:288708 3' similar to contains Alt repetitive element	
48986	17719		1.78	7.0E-01 AL168330.12	NT	Homo sapiens chromosome 21 segment HS21C101	
68862	18849		1.11	7.0E-01 AB024316.1	NT	Arabidopsis thaliana mRNA for chlorophyll b synthase, complete cds	
62276	20970		11.76	7.0E-01 AE000253.1	NT	Escherichia coli K-12 MG1655 section 143 of 400 of the complete genome	
92116	21895	35084	0.57	7.0E-01 U539888.1	NT	Clostridium acetobutylicum meroiod-specific phosphotransferase system (PTS) and mfd genes, complete cds	
92116	21895	35085	0.57	7.0E-01 U539888.1	NT	Clostridium acetobutylicum meroiod-specific phosphotransferase system (PTS) system, mfdA, mfdR, mfdF, and mfd genes, complete cds	
10526	23172	36400	0.49	7.0E-01 U34862.1	NT	Dario rufa complement factor B mRNA, complete cds	
11084	23734	37008	1.94	7.0E-01 AV763842.1	EST_HUMAN	AV763842 MDS Homo sapiens cDNA clone MDSCH-E04 5'	
11084	23734	37007	1.94	7.0E-01 AV763842.1	EST_HUMAN	AV763842 MDS Homo sapiens cDNA clone MDSCH-E04 5'	
849	13716	28380	11.02	6.9E-01 U68674.1	NT	Candida albicans equolone epoxidase (CAERG1) gene, complete cds and translatable regulator gene, partial cds	

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**Table 4**  
**Single Exon Probes Expressed In Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
949	13715	26381	11.02	6.9E-01	U68874.1	NT	<i>Candida albicans</i> squalene epoxidase (CAERG1) gene, complete cds and translational regulator gene, partial cds
1287	14037	26708	2.74	6.9E-01	AA588530.1	EST_HUMAN	mn28a09.s1 NCI_CGAP_Gen1 Homo sapiens cDNA clone IMAGE:1085176 3'
3213	15976	28827	1.97	6.9E-01	AE002271.2	NT	<i>Chlamydia muridarum</i> , section 3 of 85 of the complete genome
5694	18488	31409	0.91	6.9E-01	AB036862.1	NT	<i>Brachiolectoma belcheri</i> BbNA3 mRNA for notochord actin, complete cds
5900	18685	31683	0.82	6.9E-01	Y18278.1	NT	Drosophila melanogaster mRNA for A-Krasse anchor protein DAKAP550, partial
6277	19050	32027	1.38	6.9E-01	BE286188.1	EST_HUMAN	601177332F1 NIH_MGC_17 Homo sapiens cDNA clone IMAGE:353232B 5'
7697	20360	33474	0.65	6.9E-01	AF248883.1	NT	<i>Strongylocentrotus purpuratus</i> myosin V, complete cds
7879	20574	33700	2.98	6.9E-01	AL181573.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 69
7879	20574	33701	2.98	6.9E-01	AL181573.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 69
9069	21768		0.79	6.9E-01	AF118048.1	NT	Entamoeba dispar cation transporting ATPase (efpase) gene, partial cds
6594	22247	35431	0.59	6.9E-01	AF206319.1	NT	<i>Musa acuminate</i> pectate lyase 1 (PL1) mRNA, complete cds
6594	22247	35432	0.50	6.9E-01	AF206319.1	NT	<i>Musa acuminate</i> pectate lyase 1 (PL1) mRNA, complete cds
112223	23886	37172	2.38	6.9E-01	D89013.1	NT	Homo sapiens DAN gene, complete cds
112223	23886	37173	2.38	6.9E-01	D89013.1	NT	Homo sapiens DAN gene, complete cds
11878	25197		3.01	6.9E-01	Q89968	SWISSPROT	FORKHEAD BOX PROTEIN C2 (FORKHEAD-RELATED PROTEIN FKHL14) (MESENCHYME FORK HEAD PROTEIN 1) (MFH-1 PROTEIN) (TRANSCRIPTION FACTOR FKHL14)
887	13704	26369	1.05	6.8E-01	AF017784.1	NT	<i>Giardia intestinalis</i> cathepsin kinase gene, complete cds
2880	15339		0.99	6.8E-01	D60917.1	NT	Synechocystis sp. PCC6803 complete genome, 27/27, 3418852-3573470
2832	14358	27045	1.49	6.8E-01	AA854747.1	EST_HUMAN	6176a05_s1 Soares' parathyroid tumor NbHPA Homo sapiens cDNA clone IMAGE:1402256 3' similar to sbX56411 .mat' ALCOHOL DEHYDROGENASE CLASS II PI CHAIN (HUMAN);
4553	17268	28801	1.45	6.8E-01	J00762.1	NT	Rat(nodded) protecin gene : exon III and flanks
9538	22191	35375	1.45	6.8E-01	AB037788.1	NT	Homo sapiens mRNA for KIAA1345 protein, partial cds
11027	23689	36962	1.82	6.8E-01	AJ278876.1	NT	<i>Stagonospora avenae</i> bgf1 gene for beta-glucosidase, exons 1-4
11027	23689	36963	1.82	6.8E-01	AJ278875.1	NT	<i>Stagonospora avenae</i> bgf1 gene for beta-glucosidase, exons 1-4
11058	23728	37000	2.4	6.8E-01	AF038839.1	NT	<i>Mus musculus</i> zinc finger protein (Peg3) mRNA, complete cds
11058	23728	37001	2.4	6.8E-01	AF038839.1	NT	<i>Mus musculus</i> zinc finger protein (Peg3) mRNA, complete cds
11607	24205	37527	1.36	6.8E-01	AF110620.1	NT	Mus musculus major histocompatibility complex region NG27, NG28, RPS28, NADH oxidoreductase, NG29, KIFC1, Fes-binding protein, BING1, tapasin, RatGDS-like, KE2, BING4, beta 1,3-galactosyl transferase, and RPS18 genes, complete cds; Sacm21 gene, partial>
11607	24205	37528	1.36	6.8E-01	AF110620.1	NT	Mus musculus major histocompatibility complex region NG27, NG28, RPS28, NADH oxidoreductase, NG29, KIFC1, Fes-binding protein, BING1, tapasin, RatGDS-like, KE2, BING4, beta 1,3-galactosyl transferase, and RPS18 genes, complete cds; Sacm21 gene, partial>

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Table 4  
Single Exon Probes Expressed In Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
291	13097	25739	44.11	6.7E-01 AF213884.1	NT	Homo sapiens nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (NFKB1) gene, complete cds	
330	13131	25766	21.34	6.7E-01 AF213884.1	NT	Homo sapiens nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (NFKB1) gene, complete cds	
2143	14873	27806	1.73	6.7E-01 AA451884.1	EST_HUMAN	xx12g12.1 Soares,_lateral_fetus_Nb2H/F8_9w Homo sapiens cDNA clone IMAGE:788370 3' similar to contains element TARI repetitive element ;	
2163	16587	27628	2.51	6.7E-01 AF1886073.1	NT	Drosophila melanogaster Mst3BC gene, complete cds; NM_006470.1 (Mst3bc) gene, complete cds, alternatively spliced; and transcription factor (Relish) gene, complete cds, alternatively spliced	
2984	15780	28408	3.41	6.7E-01 6678580 NT	NT	Mus musculus Wiskott-Aldrich syndrome protein (Wasp), mRNA	
4419	17155	28786	0.79	6.7E-01 X74421.1	NT	S. tuberosum mRNA for glucose-6-phosphate dehydrogenase	
5422	18221	30832	0.94	6.7E-01 JD4838.1	NT	M.barkeri ATPase alpha and beta subunit (atpA and atpB) genes, complete cds	
5422	18221	30933	0.94	6.7E-01 JD4838.1	NT	M.barkeri ATPase alpha and beta subunit (atpA and atpB) genes, complete cds	
6231	18005	31881	1.18	6.7E-01 9835035 NT	NT	Gallid herpesvirus 2, complete genome	
6231	19075	31982	1.18	6.7E-01 9835035 NT	NT	Gallid herpesvirus 2, complete genome	
7215	19800		4.34	6.7E-01 AE004686.1	NT	Pseudomonas aeruginosa PA01, section 167 of 529 of the complete genome	
7240	18825	33000	0.92	6.7E-01 AE004686.1	NT	Halicobacter pylori, strain J89 section 47 of 132 of the complete genome	
10044	22692		0.68	6.7E-01 M34046.1	NT	Human placental protein 14 (PP14) gene, complete cds	
10873	23553	36800	2.07	6.7E-01 BES4646.1	EST_HUMAN	CA8-HT078-01080-197-c03 HT0788 Homo sapiens cDNA N-ACETYLGLUCOSAMINYL-PHOSPHATIDYLINOSITOL BIOSYNTHETIC PROTEIN GP1	
11436	23203	36435	3.59	6.7E-01 O14257	SWISSPROT		
11659	24255	37573	1.68	6.7E-01 AA342521.1	EST_HUMAN	EST148085 Fetal spleen Homo sapiens cDNA 3' end	
2505	15222	27864	1.29	6.6E-01 AF075240.1	NT	Homo sapiens SLC71 protein (SLC72) mRNA, partial cds	
2704	15411	28148	1.44	6.6E-01 AF188339.1	NT	Homo sapiens lens epithelium-derived growth factor gene, alternatively spliced, complete cds	
3650	16403	28043	4.57	6.6E-01 Y07689.1	NT	Catfishes random DNA marker, 282bp	
4099	16832		0.85	6.6E-01 UB1328.1	NT	Human hereditary haemochromatosis region, histone 2A-like protein gene, hereditary haemochromatosis (HLA-H) gene, RoRet gene, and sodium phosphatate transporter (NPT3) gene, complete cds	
5125	17843	30461	1.13	6.6E-01 AL161572.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 68	
6240	18014	31688	4.28	6.6E-01 6680577 NT	NT	Mus musculus Kinesin light chain 2 (Klc2), mRNA	
7695	20283	33359	3.76	6.6E-01 AV680508.1	EST_HUMAN	AV680508 GLC Homo sapiens cDNA clone GLCGID04 3'	
8464	21168	34289	0.52	6.6E-01 AV704700.1	EST_HUMAN	AV704700 ADB Homo sapiens cDNA clone ADBCAF1 5'	
8894	22217		2	6.6E-01 AL168278.2	NT	Homo sapiens chromosome 21 segment HS21C078	
12470	24838	31033	1.48	6.6E-01 AE004382.1	NT	Vibrio cholerae chromosome II, section 38 of 63 of the complete chromosome	
610	13388	28019	18.23	6.5E-01 M75140.1	NT	H. vulgaris Na,K-ATPase alpha subunit mRNA, complete cds	

Table 4  
Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
610	13398	26020		18.23	6.5E-01 M75140.1	NT	H. vulgaris Ns <sub>2</sub> K-ATPase alpha subunit mRNA, complete cds
3426	18183	28833		4.25	6.5E-01 AB041225.1	NT	Mus musculus gene for Tab2, complete cds
4249	16880	29816		4.23	6.5E-01 AJ272285.1	NT	Homo sapiens SPP2 gene for secreted phosphoprotein 24 precursor, exons 1-8
4277	17016	28843		0.78	6.5E-01 AL161539.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 39
5003	17728	30329		2.8	6.5E-01 U28821.1	NT	Phascolos wigeri ATPase gamma subunit mRNA, nuclear gene encoding mitochondrial protein, partial cds
5357	25067	30843		1.77	6.5E-01 P18480	SWISSPROT	TRANSCRIPTION REGULATORY PROTEIN SNFs (SWISSNFCOMPLEX COMPONENT SNF5) (TRANSCRIPTION FACTOR TFE4)
6827	18424	31337		0.62	6.5E-01 AL163249.2	NT	Homo sapiens chromosome 21 segment HS21C049
6825	19387	32400		1.5	6.5E-01 D88348.1	NT	Chicken mRNA for 115-kDa melanosome matrix protein, complete cds
7668	20236	33340		0.84	6.5E-01 AI788882.1	EST_HUMAN	wc46802.x1 NCI_CGAP_P728 Homo sapiens cDNA clone IMAGE:23216423'
9737	22388			0.8	6.5E-01 T78904.1	EST_HUMAN	yd21b04.s1 Scores fetal liver spleen TINF2 Homo sapiens cDNA clone IMAGE:1088473'
10233	22881	36084		1.98	6.5E-01 AF19878.1	NT	Mus musculus small GTP-binding protein RAB25 (Rab25) gene, complete cds
10629	23226	36460		2.68	6.5E-01 H87583.1	EST_HUMAN	yw1706.s1 Scores_plecentia_8to9weeks_21bHP8tbsW Homo sapiens cDNA clone IMAGE:2625165'
10585	23280	36518		3.5	6.5E-01 AA801287.1	EST_HUMAN	nt!bc7.5t1 NCI_CGAP_Phe1 Homo sapiens cDNA clone IMAGE:1100748.3'
10590	23381			3.83	6.5E-01 AU138078.1	EST_HUMAN	AU138078 PLACE1 Homo sapiens cDNA clone PLACE107810.5'
111598	24198	37518		2.42	6.5E-01 AF014115.1	NT	Plasmiodium berghei cytochrome c oxidase subunit III, cytochrome c oxidase subunit I, and cytochrome b genes, mitochondrial proteins, complete cds
12287	24710			2.07	6.6E-01 BE465050.1	EST_HUMAN	hn74a10x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3179130.3'
12504	25146			1.81	6.5E-01 Z74-45.1	NT	S.cerevisiae chromosome IV reading frame ORF YLD97c
245	13054	25694		8.05	6.4E-01 U48848.1	NT	Drosophila melanogaster 8kd dynamin light chain mRNA, complete cds
2803	15307	28043		1.18	6.4E-01 AF161184.1	NT	Pseudomonas fluorescens tryptophan halogenase (pmA) gene, complete cds
3449	16205	28855		2.18	6.4E-01 U48884.2	NT	Mus musculus dystroglycan 1 (DAG1) gene, exons 1 and 2 and complete cds
38442	16893	29230		1.08	6.4E-01 AB046827.1	NT	Homo sapiens mRNA for KIAA1607 protein, partial cds
8510	21202	34347		1.82	6.4E-01 AE001247.1	NT	Treponema pallidum section 63 of 87 of the complete genome
8889	22637	35848		8.8	6.4E-01 U82828.1	NT	Homo sapiens ataxia telangiectasia (ATM) gene, complete cds
10004	22652	35884		1.22	6.4E-01 BF870405.1	EST_HUMAN	602150289f1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4291128.6'
12382	24777			5.89	6.4E-01 AV7789212.1	EST_HUMAN	AV7789212 MDS Homo sapiens cDNA clone MDSCGC09.6'
425	13211	25898		4.58	6.3E-01 Po5228	SWISSPROT	HISTIDINE-RICH PROTEIN PRECURSOR (CLONE PFHRP-II)
522	13306	25938		2.25	6.3E-01 U32689.1	NT	Haemophilus influenzae Rd section 4 of 163 of the complete genome
2159	14889	27623		2.02	6.3E-01 U81136.1	NT	Shigella flexneri multi-antibiotic resistance locus
2553	15297	28035		3.51	6.3E-01 U75331.1	NT	Gallus gallus bone morphogenic protein 1 (BMP1) mRNA, partial cds
2553	15297	28036		3.51	6.3E-01 U75331.1	NT	Gallus gallus bone morphogenic protein 1 (BMP1) mRNA, partial cds

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**Table 4**  
**Single Exon Probes Expressed In Brain**

Probe Seq ID No:	Exon Seq ID No:	ORF Seq ID No:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
6973	18755	31716	0.94	6.3E-01	BE5083988.1	EST_HUMAN	PM0-BT0757-010500-002-005 BT0757 Homo sapiens cDNA
6504	19269	32271	0.84	6.3E-01	L277888.1	NT	Streptococcus dysgalactiae (mag) gene, complete cds
6504	19269	32272	0.84	6.3E-01	L277888.1	NT	Streptococcus dysgalactiae (mag) gene, complete cds
8419	21112		3.44	6.3E-01	BE502044.1	EST_HUMAN	601676889F1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3959351 6'
8784	21478	34624	0.95	6.3E-01	S82927.1	NT	glycoprotein IIIa (Alu 1 and 3 fusion junction) [human, Genomic Mutant, 300 n]
9120	21808	34975	0.8	6.3E-01	BF216884.1	EST_HUMAN	6018BA050F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:41025688 5'
9320	21887	35159	2.45	6.3E-01	9627521	NT	Variola virus, complete genome
9320	21887	35160	2.45	6.3E-01	9627521	NT	Variola virus, complete genome
98338	22489		0.67	6.3E-01	AE02328.2	NT	Chlamydia muridarum, section 68 of 85 of the complete genome
10328	22873	36183	1.47	6.3E-01	Z73003.1	NT	S.cerevisiae chromosome VII reading frame ORF YGR218w
10427	23073	36294	1.19	6.3E-01	AE000313.1	NT	Escherichia coli K-12 MG1655 section 203 of 400 of the complete genome
10456	23102		0.45	6.3E-01	AW785385.1	EST_HUMAN	PM0-UM0108-130500-003-012 UM0108 Homo sapiens cDNA
10983	23867	36924	2.21	6.3E-01	AA877715.1	EST_HUMAN	IT0906.s1 NCI_CGAP_Colo Homo sapiens cDNA clone IMAGE:1161371 3' similar to TR:002816 002816
11308	23867	37268	9.25	6.3E-01	AI904160.1	EST_HUMAN	CHM-BT043-080289-048 BT043 Homo sapiens cDNA
11402	24051	37355	1.68	6.3E-01	P47003	SWISSPROT	HYPOTHETICAL 13.7 KD PROTEIN IN INO1-DS2 INTERGENIC REGION
11681	24180	37495	1.84	6.3E-01	P36073	SWISSPROT	HYPOTHETICAL 16.3 KD PROTEIN IN VWA12-APN1 INTERGENIC REGION
11988	25355	30607	4.37	6.3E-01	8910283	NT	Mus musculus keratin complex 2, gene 8g (Krt2-8g), mRNA
12078	24587		1.45	6.3E-01	AF105227.1	NT	Homo sapiens 3'-phosphoadenosine 5'-phosphosulfate synthetase (PAPS) mRNA, complete cds
12283	25272		2.93	6.3E-01	X83628.1	NT	Clinicode pcd gene
5780	18571	31489	2.31	6.2E-01	Q10135	SWISSPROT	HYPOTHETICAL 142.5 KD PROTEIN C235E2.02 IN CHROMOSOME 1
73394	20073		3.44	6.2E-01	AF022283.1	NT	Mus musculus calcium-sensing receptor related protein 4 (Casr-r4) mRNA, partial cds
7443	25114	33209	1.33	6.2E-01	AL021127.2	NT	Mus musculus chaperone X antigen A, putative Mage8 gene, Caiitactin, NAD(P) steroid dehydrogenase and Zinc finger protein 185
8200	20894	34031	4.52	6.2E-01	H72285.1	EST_HUMAN	ys01608.s1 Severe fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:2135423
6755	21447	34525	0.52	6.2E-01	AF034411.1	NT	Lycopersicon esculentum cytochrome Cu,Zn superoxide dismutase (Sod) gene, partial cds; end dehydroquinat
9349	20420	33540	1.55	6.2E-01	BE562687.1	EST_HUMAN	dehydrazase/shiftin-like/NADP oxidoreductase gene, complete cds
8410	22072		2.55	6.2E-01	M24461.1	NT	Human pulmonary surfactant-associated protein SP-B (SPFB3) mRNA, complete cds
8978	22826	35834	6.2	6.2E-01	AL161511.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 23
10121	22769	35982	0.5	6.2E-01	11420793	NT	Homo sapiens potassium voltage-gated channel, Shab-related subfamily, member 1 (KCNB1), mRNA
10121	22769	35983	0.5	6.2E-01	11420793	NT	Homo sapiens potassium voltage-gated channel, Shab-related subfamily, member 1 (KCNB1), mRNA

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**Table 4**  
**Single Exon Probes Expressed in Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
10434	23080	36305		5.2	6.2E-01 P27410	SWISSPROT	NON-STRUCTURAL POLYPROTEIN [CONTAINS: RNA-DIRECTED RNA POLYMERASE ; THIOL PROTEASE PAC ; HELICASE (2C LIKE PROTEIN); COAT PROTEIN]
10434	23080	36308		5.2	6.2E-01 P27410	SWISSPROT	NON-STRUCTURAL POLYPROTEIN [CONTAINS: RNA-DIRECTED RNA POLYMERASE ; THIOL PROTEASE P3C ; HELICASE (2C LIKE PROTEIN); COAT PROTEIN]
2283	15114			4.38	6.1E-01 6578076 NT	Mus musculus secreted acidic cysteine rich glycoprotein (Sparc), mRNA	
6449	18248	31137		1.15	6.1E-01 M58940.1	Caenorhabditis elegans N2 CetMycD (hlh-1) alternatively spliced genes, complete cds	
6770	19514	32540		4.02	6.1E-01 M84733.1	Rat TRPM-2 gene, complete cds	
6770	19514	32541		4.02	6.1E-01 M84733.1	Rat TRPM-2 gene, complete cds	
6820	18856	32702		0.84	6.1E-01 AW105833.1	EST_HUMAN	X50103_X1 NC1 CGAP_Ov23 Homo sapiens cDNA clone IMAGE:2597237 3' similar to dbX12671_m81
7005	18697	32751		0.72	6.1E-01 Q63768	SWISSPROT	HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN A1 (HUMAN); SUSHI REPEAT-CONTAINING PROTEIN SRPX PRECURSOR (DRS PROTEIN) (DOWN-REGULATED BY V-SRO)
8132	20826	33962		3.27	6.1E-01 AF038353.1	NT	Arabidopsis thaliana putative zinc transporter (ZIP1) mRNA, complete cds
8694	21388	34528		1.09	6.1E-01 11431055 NT	Homo sapiens mitogen-activated protein kinase kinase kinase 4 (MAP4K4), mRNA	
8694	21388	34529		1.09	6.1E-01 11431065 NT	Homo sapiens mitogen-activated protein kinase kinase kinase 4 (MAP4K4), mRNA	
9315	21982	35153		18.74	6.1E-01 AF238617.1	NT	Homo sapiens G-protein coupled receptor EDG-7 mRNA, complete cds
9315	21982	35154		18.74	6.1E-01 AF238617.1	NT	Homo sapiens G-protein coupled receptor EDG-7 mRNA, complete cds
9742	22383	35697		0.93	6.1E-01 AE004452.1	NT	Pseudomonas aeruginosa PA01, section 13 of 528 of the complete genome
8946	22594	35797		1.06	6.1E-01 AF119117.1	NT	Homo sapiens dopamine transporter (SLC6A3) gene, complete cds
11738	24331	37655		2.57	6.1E-01 SB3182.1	NT	heterogeneous-binding protein-hepatocyte growth factor activator homolog [human, plasma, mRNA, 2408 nt]
11738	24331	37656		2.57	6.1E-01 SB3182.1	NT	heterogeneous-binding protein-hepatocyte growth factor activator homolog [human, plasma, mRNA, 2408 nt]
12074	25159	30889		2.28	6.1E-01 AB041350.1	NT	Mus musculus Cdh45 mRNA for type IV collagen alpha 5 chain, complete cds
12694	24977			1.57	6.1E-01 X95287.1	NT	M.musculus Cdh45 mRNA for type IV collagen alpha 5 chain, complete cds
482	13287	25693		1.24	6.0E-01 D87875.1	NT	Homo sapiens DNA for amyloid precursor protein, complete cds
548	13331			3.09	6.0E-01 5802899 NT	Human respiratory syncytial virus strain CLA20, mRNA	
1341	14089	26765		1.91	6.0E-01 AF065293.1	NT	Human respiratory syncytial virus strain CH93-53b attachment protein (G) gene, complete cds
3795	16547	29180		0.92	6.0E-01 AJ233396.1	NT	Viral hemophagic septicemia virus N, P, M, G, NV, L genes, French strain 07-7-1
4165	16905			1.09	6.0E-01 AF058896.1	NT	Homo sapiens Notch3 (NOTCH3) gene, exons 28, 27, and 28
5189	18007	30828		2	6.0E-01 P20288	SWISSPROT	D(2) DOPAMINE RECEPTOR
5333	18166	30839		2.86	6.0E-01 AW139773.1	EST_HUMAN	U-H-B11-acb-a-10-0-U.1st NC1_CGAP_Sub3 Homo sapiens cDNA clone IMAGE:27188193'
6445	19213	32210		2.78	6.0E-01 U38813.1	NT	Musca domestica insecticide-susceptible strain voltage-sensitive sodium channel mRNA, complete cds

**Table 4**  
**Single Exon Probes Expressed In Brain**

Probe SEQ ID NO:	Exon seq ID NO:	ORF seq ID NO:	Expression Signal	Most Similar BLAST E Value	Top Hit Accession No.	Top Hit Database Sources	Top Hit Descriptor
6563	19328	32335	0.68	6.0E-01	Q04912	SWISSPROT	MACROPHAGE-STIMULATING PROTEIN RECEPTOR PRECURSOR (MSP RECEPTOR) (P168-RON) (CDW138) (CD138 ANTIGEN)
7254	19538	33013	6.98	6.0E-01	AL277681.1	NT	Homo sapiens partial LMO1 gene for LIM domain only 1 protein, exon 1
8023	20718	33850	4.39	6.0E-01	P02835	SWISSPROT	SEGMENTATION PROTEIN FUSHI TARAZU
8923	20718	33851	4.39	6.0E-01	P02835	SWISSPROT	SEGMENTATION PROTEIN FUSHI TARAZU
9723	22374	35574	1.61	6.0E-01	AB0081183.1	NT	Homo sapiens genes for leukotriene B4 receptor BLT2, leukotriene B4 receptor BLT1, complete cds
10173	22821		1.46	6.0E-01	Q01497	SWISSPROT	PEROXISOMAL MEMBRANE PROTEIN PER9 (PEROXIN-3)
10980	23664	36921	1.49	6.0E-01	AJ131892.1	NT	Ceinus gallicus mRNA for Hyperian protein, 419 kD isoform
10980	23664	36922	1.49	6.0E-01	AJ131892.1	NT	Ceinus gallicus mRNA for Hyperian protein, 419 kD isoform
11640	24140	37449	3.77	6.0E-01	AJ420623.1	EST_HUMAN	fl0807_x1 NCI_CGAP_Pt28 Homo sapiens cDNA clone IMAGE:2095821 3'
12354	24768	31050	2.25	6.0E-01	11421683	NT	Homo sapiens nuclear factor erythroid-derived 2-like 3 (NFE2L3), mRNA
12455	24824		2.6	6.0E-01	AA706037.1	EST_HUMAN	Z98g05_31 Soenes_fetal_fov_spleen_cDNA clone IMAGE:4627783
12639	252038	30815	3.04	6.0E-01	9055303	NT	Mus musculus cGMP-inhibited phosphodiesterase (Pde3a), mRNA
12639	25142		2.03	6.0E-01	BE167617.1	EST_HUMAN	RC1-HT0375-030500-016-c03 HT0376 Homo sapiens cDNA
980	13745	28407	1.38	6.9E-01	UR22701.1	NT	Haemophilus influenzae Rd section 16 of 163 of the complete genome
3264	16028	28675	2.29	6.9E-01	AL163267.2	NT	Homo sapiens chromosome 21 segment HS21C087
3264	16026	28676	2.29	6.9E-01	AL163267.2	NT	Homo sapiens chromosome 21 segment HS21C087
4198	16937		4.21	6.9E-01	AF162766.1	NT	Rettis norvegicus cerebellum 2 mRNA, partial cds
6973	19142	32139	1.56	5.9E-01	AF085440.2	NT	Homo sapiens low density lipoprotein receptor-related protein II (LRP2) gene, exon 1 and partial cds
7166	19853	32822	1.32	5.9E-01	AB022488.1	NT	Homo sapiens gene for histamine H2 receptor, promoter region and complete cds
7286	19979		0.61	5.9E-01	X68801.1	NT	G. gallus gene for skeletal alpha-actinin, exon EF2
7898	20593	33725	0.48	5.9E-01	D80911.1	NT	Synecocystis sp. PCC6803 complete genome, 13227, 1576893-1719843
8536	21228	34370	0.48	5.9E-01	D12822.1	NT	Legionella pneumophila gene for iron superoxide dismutase, complete cds
9443	22120	35289	0.82	5.9E-01	AF083204.2	NT	Chlamydia trachomatis strain KUW31/Cx major outer membrane protein (omp1) gene, complete cds
98113	22464		0.74	5.9E-01	P08483	SWISSPROT	E6 PROTEIN
10088	22738	35951	1.15	5.9E-01	P66284	SWISSPROT	VASCULAR ENDOTHELIAL-CADHERIN PRECURSOR (VE-CADHERIN) (CADHERIN-5)
10569	23284	36502	2.5	5.9E-01	Q8X013	SWISSPROT	THYMIDYLATE KINASE (DTMP KINASE)
10576	23271	36507	1.72	5.9E-01	AF197044.1	NT	Xenopus laevis receptor protein tyrosine phosphatase delta (XPTP-D) mRNA, complete cds
10881	23561	36808	2.91	5.9E-01	AW837175.1	EST_HUMAN	PMI-DT0041-180100-002-h03 DT0041 Homo sapiens cDNA
11149	23816	37098	1.95	5.9E-01	AF064623.1	NT	Mus sprattus strath SPRETIEI CD48 antigen (Cd48) gene, partial cds
11458	24062	37358	1.55	5.9E-01	P47135	SWISSPROT	JSN1 PROTEIN

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Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
11458	24062	37359	1.58	5.8E-01	P47135	SWISSPROT	J3N1 PROTEIN
12021	24549	31109	2	5.8E-01	LA2320_1	NT	Oryctolagus cuniculus alpha 1 anti-trypsin (alpha 1 AT) gene, promoter region
12252	24898		4.35	5.8E-01	AB017706_1	NT	Aspergillus oryzae pyrG gene for orotidine-5'-phosphate decarboxylase, complete cds
12465	24832		5.72	5.8E-01	P34928	SWISSPROT	MICROTUBULE-ASSOCIATED PROTEIN 1A [CONTAINS: MAP1 LIGHT CHAIN [LC2]]
18022	14639	27348	1.36	5.8E-01	P40472	SWISSPROT	SIM1 PROTEIN
25599	16283	28021	1.01	5.8E-01	7305230	NT	<i>Mus musculus</i> low-density lipoprotein B (ldlb), mRNA
4478	17213	28838	4.37	5.8E-01	AB009077_1	NT	Vigma radiata mRNA for proton pyrophosphatase, complete cds
52290	18095		0.82	5.8E-01	AED02162_1	NT	Ureaplasma urealyticum section 53 of 59 of the complete genome
54444	18243	31131	0.62	5.8E-01	Q10659	SWISSPROT	POTENTIAL 5'-3' EXONUCLEASE
5991	18869	31835	1.09	5.8E-01	D78658_1	EST_HUMAN	HUM500E068 Human placenta polyA+ (TF44mRNA) Homo sapiens cDNA clone GEN-500E068
62220	18994	31970	0.66	5.8E-01	D50601_1	NT	<i>Shigella sonnei</i> DNA for 28 ORFs, complete cds
67115	19830		2.48	5.8E-01	S65091_1	NT	cyclic AMP-regulated phosphoprotein [rats, mRNA, 1030 nt]
7787	20482		2.61	5.8E-01	I441571_1	EST_HUMAN	<i>yne1</i> [C. elegans] Soanes adult brain N2b; FB55Y Homo sapiens cDNA clone IMAGE:175757 3' similar to
7885	20680	33805	0.64	5.8E-01	AI28005_1	EST_HUMAN	gb:S78187 M-PHASE INDUCER PHOSPHATASE 2 (HUMAN); qhs5d10_xt_Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1853779 3'
7885	20680	33806	0.64	5.8E-01	AI28005_1	EST_HUMAN	qhs5d10_xt_Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1853779 3'
8080	20784	33914	3.41	5.8E-01	P14328	SWISSPROT	SPORE COAT PROTEIN SP88
8090	20784	33915	3.41	5.8E-01	P14328	SWISSPROT	SPORE COAT PROTEIN SP88
8789	21481	34628	8.87	5.8E-01	AJ270774_1	NT	<i>Homo sapiens</i> partial TCF-4 gene for T-cell transcription factor-4, exons 6-11
8871	21562	34707	0.98	5.8E-01	Q27368	SWISSPROT	TRANSCRIPTION FACTOR E2F
8872	21563	34708	0.51	5.8E-01	Q20471	SWISSPROT	PUTATIVE CASEIN KINASE I F46F22 IN CHROMOSOME X
9496	22149		0.81	5.8E-01	BF031688_1	EST_HUMAN	60155774f1 NIH_MGC_58 Homo sapiens cDNA clone IMAGE:3827288 5'
10811	23391	36837	7.58	5.8E-01	AJ243213_1	NT	<i>Homo sapiens</i> partial 5-HT4 receptor gene, exons 2 to 5
10862	23638		3.97	5.8E-01	BF700092_1	EST_HUMAN	602127677f1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4284403 5'
11089	23769		1.99	5.8E-01	BF700082_1	EST_HUMAN	602127577f1 NIH_MGC_56 Homo sapiens cDNA clone IMAGE:4284403 5'
1480	14227	26912	1.12	5.7E-01	P08727	SWISSPROT	APOLIPOPROTEIN A-IV PRECURSOR (APO-AIV)
1480	14227	26913	1.12	5.7E-01	P08727	SWISSPROT	APOLIPOPROTEIN A-IV PRECURSOR (APO-AIV)
3033	15804		0.69	5.7E-01	6755253	NT	<i>Mus musculus</i> plasmacytoma variant translocation 1 (Pvt1), mRNA
3217	15980	28831	1.62	5.7E-01	Q9WTJ2	SWISSPROT	PUTATIVE TRANSCRIPTION FACTOR OVO-LIKE 1 (MOVO1A)
3485	16251		2.82	5.7E-01	AB033563_1	NT	<i>Populus tremuloides</i> pectic-2 mRNA for 1-aminocyclopentane-1-carboxylate synthase, complete cds
6262	18036	32011	5.13	5.7E-01	BF036413_1	EST_HUMAN	6014549622f1 NIH_MGC_88 Homo sapiens cDNA clone IMAGE:3858590 5'
6811	18374	32388	0.81	5.7E-01	AA184201_1	EST_HUMAN	323806.f1 Soanes_NIH-MPU_S1 Homo sapiens cDNA clone IMAGE:665674 5'
6763	17832	30568	1.33	5.7E-01	AL111440_1	NT	<i>Batrachochytrium degenerei</i> strain T4 cDNA library under conditions of nitrogen deprivation

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**Table 4**  
**Single Exon Probes Expressed In Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
7684	20328	33438	2.14	5.7E-01	P00373	SWISSPROT	PYRROLINE-5-CARBOXYLATE REDUCTASE (P5CRL) (P5CREDUCTASE)
7870	20665		0.5	6.7E-01	AA1251835.1	NT	Mus musculus Kcnq1 / Lmpc5, Mash2, Tapa-1, Tsc2 and Tsc2b genes, alternative transcripts
8279	20973		0.47	6.7E-01	AI086061.1	EST_HUMAN	HA0585 Human fetal liver cDNA library Homo sapiens cDNA
8699	22350	35544	1.19	6.7E-01	AL1618322	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 32
8699	22350	35545	1.19	6.7E-01	AL1618322	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 32
10475	23121	36351	0.72	6.7E-01	BF540982.1	EST_HUMAN	602087712F1 NIH MGIC_68 Homo sapiens cDNA clone IMAGE:4088610 5
11983	24524		1.92	6.7E-01	BE715051.1	EST_HUMAN	MR3-HT0738-180700-003-ed2 HT0738 Homo sapiens cDNA
12658	24958		3.01	5.7E-01	BE869722.2	EST_HUMAN	601654814R1 NIH MGIC_57 Homo sapiens cDNA clone IMAGE:3839783 3
3357	18117	28772	1.3	6.8E-01	AB018283.2	NT	Homo sapiens mRNA for KIAA0740 protein, partial cds
3357	18117	28773	1.3	5.6E-01	AB018283.2	NT	Homo sapiens mRNA for KIAA0740 protein, partial cds
3883	16813	29252	0.97	5.6E-01	AI161601.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 13
4215	16953	29578	0.74	5.6E-01	D83136.1	NT	Chicken TBP gene, exon8, complete cds
8702	21394	34541	4.01	5.6E-01	AV684703.1	EST_HUMAN	AV684703 GKC Homo sapiens cDNA clone GKCCSF05 5
8702	21394	34542	4.01	5.6E-01	AV684703.1	EST_HUMAN	AV684703 GKC Homo sapiens cDNA clone GKCCSF05 5
9275	22029	35189	1.08	5.6E-01	AB0388782.1	NT	Homo sapiens MUC5A gene for intestinal mucus, partial cds
11884	24467		2.57	5.6E-01	BE888280.1	EST_HUMAN	601514007F1 NIH MGIC_71 Homo sapiens cDNA clone IMAGE:3815457 5
11897	24535	37272	1.63	5.6E-01	AA483535.1	EST_HUMAN	ng75q10.s1 NCI CGAP_P16 Homo sapiens cDNA clone IMAGE:940674 similar to contains element P177 repetitive element;
122352	18613	28252	1.69	6.6E-01	AL161501.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 13
12379	24776		2.7	5.6E-01	PS0505	SWISSPROT	HIGH AFFINITY POTASSIUM TRANSPORTER
12773	25027		4.23	5.6E-01	BFF7328.1	EST_HUMAN	602132024F1 NIH MGIC_81 Homo sapiens cDNA clone IMAGE:4271334 5
1189	13941	28606	0.85	6.6E-01	83883912	NT	Rattus norvegicus Propriety Coenzyme A carboxylase, beta polypeptide (Pob), mRNA
2705	15412	28149	6.93	5.5E-01	P03341	SWISSPROT	GAG POLYPROTEIN [CONTAINS: INNER COAT PROTEIN P12; CORE PROTEIN P15; CORE SHELL PROTEIN P30; NUCLEOPROTEIN P10]
2705	18412	28160	6.98	5.5E-01	P03341	SWISSPROT	GAG POLYPROTEIN [CONTAINS: INNER COAT PROTEIN P12; CORE PROTEIN P15; CORE SHELL PROTEIN P30; NUCLEOPROTEIN P10]
2819	16685	28330	1	5.5E-01	690202085	NT	Homo sapiens superkiller viralicidic activity 2 (S. cerevisiae homolog)-like (SK172), mRNA
3082	15823		1.55	5.5E-01	H46218.1	EST_HUMAN	y01Ba10.s1 Soares adult brain N268-1865Y Homo sapiens cDNA clone IMAGE:176268 3
3228	15891	28844	4.22	5.5E-01	AF227240.1	NT	Rabbit oral papillomatitus, complete genome
3678	18431	28073	1.7	5.5E-01	P48765	SWISSPROT	FOS-RELATED ANTIGEN-1
5082	17801	30419	1.79	5.5E-01	U69397.1	NT	Bos taurus NHC class II Beta-chain Bta-L-DIB1 gene, partial cds
7187	18873		0.66	5.5E-01	AB015598.1	NT	Ceratostis auratus gene for gonadotroph II beta subunit, complete cds
8348	21041	34178	1.04	5.5E-01	AI791768.1	EST_HUMAN	crd2c01.s1 NCI CGAP_Lu5 Homo sapiens cDNA clone IMAGE:6022336 5
9887	22349		0.7	5.5E-01	U88415.1	NT	Crimean-Congo hemorrhagic fever virus strain SPU 41/G85 nucleoprotein gene, complete cds

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Table 4  
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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
10276	22824	36138	0.98	5.6E-01	T05047.1	EST_HUMAN	EST02835 Fetal brain, Strategene (cat#336206) Homo sapiens cDNA clone HFBCQ35
11087	23757	37033	1.65	5.6E-01	BF128507.1	EST_HUMAN	601811077R1 NIH_MGC_48 Homo sapiens cDNA clone IMAGE-4054003 3'
140	12955	25597	4.91	5.4E-01	7657288 NT		Homo sapiens KIAA0929 protein Msx2 interacting nuclear target (MINT) homolog (KIAA0929), mRNA
140	12955	25598	4.91	5.4E-01	7657288 NT		Homo sapiens KIAA0929 protein Msx2 interacting nuclear target (MINT) homolog (KIAA0929), mRNA
571	13352	25980	1.18	5.4E-01	AF232008.1	NT	Pseudomonas syringae pv. tomato strain DC3000 AvrE (avrE), HrpW (hrpW), and GstA (gstA) genes, complete cds; and unknown genes
571	13352	25981	1.18	5.4E-01	AF232008.1	NT	Pseudomonas syringae pv. tomato strain DC3000 AvrE (avrE), HrpW (hrpW), and GstA (gstA) genes, complete cds; and unknown genes
1248	13987	26884	3.41	5.4E-01	AW886087.1	EST_HUMAN	QV4-NR0040-070400-1160-c04 NH0040 Homo sapiens cDNA
2089	14830		3.43	5.4E-01	AE002247.2	NT	Chlamydomonas pneumoniasis AR39, section 74 of 84 of the complete genome
2252	14980	27718	1.91	5.4E-01	AJ276882.1	NT	Drosophila melanogaster mRNA for 11,15' beta carotene dioxygenase (beta-diox gene)
5056	17785	30402	0.92	5.4E-01	M74439.1	NT	Rattus norvegicus UDP glucose:xylose transferase gene, complete cds
6571	18389	31278	0.74	5.4E-01	AW842327.1	EST_HUMAN	PM2-CN1030-030200-003-c10 CN1030 Homo sapiens cDNA
6098	18876	31845	0.38	5.4E-01	AB025017.1	NT	Rattus norvegicus gene for TIS11, complete cds
6828	19684	32710	0.97	5.4E-01	BE9868682.2	EST_HUMAN	601880278R1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE-39068080 3'
7235	18920	32893	0.91	5.4E-01	221618.1	NT	S.cerevisiae RIB3 gene encoding DBP synthase
7235	18920	32894	0.91	5.4E-01	221618.1	NT	S.cerevisiae RIB3 gene encoding DBP synthase
							MITOCHONDRIAL TRIFUNCTIONAL ENZYME ALPHA SUBUNIT PRECURSOR (TP-ALPHA) [INCLUDES: LONG-CHAIN ENOYL-COA HYDRATASE ; LONG CHAIN 3-HYDROXYACYL-COA DEHYDROGENASE ]
7237	18922	32897	1.48	5.4E-01	Q84428	SWISSPROT	602076845F1 NIH_MGC_62 Homo sapiens cDNA clone IMAGE-4243680 5'
88890	22540		2.08	5.4E-01	BF572836.1	EST_HUMAN	NITRATE REDUCTASE [NADPH] (NR)
11016	23687	36948	2.87	5.4E-01	P36858	SWISSPROT	LAMININ ALPHA-2 CHAIN PRECURSOR (LAMININ M CHAIN) (MEROSIN HEAVY CHAIN).
11621	24218	37541	3.08	5.4E-01	Q80675	SWISSPROT	LAMININ ALPHA-2 CHAIN PRECURSOR (LAMININ M CHAIN) (MEROSIN HEAVY CHAIN)
11644	24489		3.5	5.4E-01	Q604X1 NCI_CGAP_U11 Homo sapiens cDNA clone IMAGE-2427128 3' similar to g:M13452 LAMIN A (HUMAN);		
603	13287	25821	1.54	5.3E-01	AF019413.1	NT	Homo sapiens HLA class III region containing tenascin X (tenascin-X) gene, partial cds; cytochrome P450 21-Hydroxylase (CYP21B), complement component C4 (C4B) G11, helicase (SKI2W), RD, complement factor B (BF), and complement component C2 (C2) genes. >
2136	14866	27598	1.01	5.3E-01	AF113919.1	NT	Brassica oleracea var. capitata phospholipase D2 (PLD2) gene, complete cds
2136	14868	27597	1.01	5.3E-01	AF113919.1	NT	Brassica oleracea var. capitata phospholipase D2 (PLD2) gene, complete cds
2786	15491	28230	6.83	5.3E-01	4506328 NT		Homo sapiens protein tyrosine phosphatases, receptor-type, zeta polypeptide 1 (PTPRZ1) mRNA

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Table 4  
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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar BLAST E Value	(Top) Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
2788	15491	28231	6.83	5.3E-01	4506328 NT	Homo sapiens protein tyrosine phosphatase, receptor-type, zeta polypeptide 1 (PTPRZ1) mRNA	
3237	15898	28349	2.74	5.3E-01	AF087658.1 NT	Homo sapiens secreted C-type lectin precursor (LSLCL) gene, complete cds	
4187	16828		1.98	5.3E-01	U39687.1 NT	Mycoplasma genitalium section 9 of 51 of the complete genome	
5371	18172	30960	1.98	5.3E-01	AI620821.1 EST HUMAN	ZM42h12.5 Scores over tumor NbHOT Homo sapiens cDNA clone IMAGE:7407115'	
5371	18172	30981	1.98	5.3E-01	AI620821.1 EST HUMAN	ZM42h12.5 Scores over tumor NbHOT Homo sapiens cDNA clone IMAGE:7407115'	
5468	18285	31156	0.94	5.3E-01	AA183672.1 EST HUMAN	ZM42g09.1 Scores NihMFu_S1 Homo sapiens cDNA clone IMAGE:6681125'	
5468	18285	31157	0.94	5.3E-01	AA183672.1 EST HUMAN	ZM42g09.1 Scores NihMFu_S1 Homo sapiens cDNA clone IMAGE:6681125'	
5559	18358	31266	1.82	5.3E-01	BE8945820.1 EST HUMAN	ZM73g12.x1 NCI_CGAP_Pz28 Homo sapiens cDNA clone IMAGE:3288118 3' similar to gb:J02783	
65559	18358	31287	1.82	5.3E-01	BE8945820.1 EST HUMAN	ZM73g12.x1 NCI_CGAP_Pz28 Homo sapiens cDNA clone IMAGE:3288118 3' similar to gb:J02783	
8802	21484		1.8	6.3E-01	LO1980.2 NT	Ranidula gorgonias fibulosa 1,5-bisphosphate carboxylase (rbcl.) gene, partial cds; chloroplast gene for chloroplast product	
8854	21545	34692	0.81	5.3E-01	BF433956.1 EST HUMAN	ZM771c12.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3 3' similar to contains element MER29	
8854	21545	34693	0.81	5.3E-01	BF433956.1 EST HUMAN	ZM771c12.x1 NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE:3 3' similar to contains element MER29	
10111	22759	35971	0.82	5.3E-01	AI654210.1 EST HUMAN	WY94b22.x1 NCI_CGAP_M15 Homo sapiens cDNA clone IMAGE:2351276 3' similar to SW:COXA_HUMAN P20674 CYTOCHROME C OXIDASE POLYPEPTIDE VA PRECURSOR;	
11550	24149	37460	7.3	5.3E-01	BE566229.1 EST HUMAN	6013398677f1 NIH MGC_63 Homo sapiens cDNA clone IMAGE:3882168 6'	
11789	24379	37709	1.72	5.3E-01	Q05783 SWISSPROT	BASEMENT MEMBRANE-SPECIFIC HEPARAN SULFATE PROTEOGLYCAN CORE PROTEIN PRECURSOR (HSPG) (PERLECAN) (PLC)	
11877	25206		4.03	5.3E-01	AA916033.1 EST HUMAN	Q930d5.51 NCI_CGAP_B77 Homo sapiens cDNA clone IMAGE:1441376 3' similar to gb:J02811	
797	13569	26229	18.35	5.2E-01	L20770.1 NT	APOLIPOPROTEIN D PRECURSOR (HUMAN); Drosophila melanogaster helicase-loop-helicase, complete cds	
11441	13896	28557	8.29	5.2E-01	Q9WV30 SWISSPROT	NUCLEAR FACTOR OF ACTIVATED T CELLS 6 (T CELL TRANSCRIPTION FACTOR NFAT5) (NFAT5) (REL DOMAIN-CONTAINING TRANSCRIPTION FACTOR NFAT5)	
1169	13923	28585	1.77	5.2E-01	AF224492.1 NT	Homo sapiens phosphatidylserine 1 gene, complete cds	
1879	14816		2.35	5.2E-01	AL163285.2 NT	Homo sapiens chromosome 21 segment HS21C085	
2142	14872	27605	2.55	5.2E-01	AB018283.2 NT	Homo sapiens mRNA for KIAA0740 protein, partial cds	
3117	15882	28521	1.23	5.2E-01	U65842.1 NT	Chlamydomonas abortus strain S28/S POM/PegA and POMP90A precursor, genes, complete cds	
3231	15994		1	5.2E-01	D73443.1 NT	Azotobacter vinelandii lsd gene for lactate dehydrogenase, complete cds	
3400	16158		1.58	5.2E-01	AL116780.1 NT	Bordetella cinesiae strain T4 cDNA library under conditions of nitrogen deprivation	
3437	16163	28643	2.27	5.2E-01	AA984165.1 EST HUMAN	em77g05.51 Strategene schizo brain S11 Homo sapiens cDNA clone IMAGE:1616504 3'	

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**Table 4**  
**Single Exon Probes Expressed In Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
3623	16378		0.76	5.2E-01	AF020269.1	NT	Medicago sativa chloroplast malaes dehydrogenase precursor (p1meth) mRNA, nuclear gene encoding chloroplast protein, complete cds
45688	17303	28530	0.82	5.2E-01	6752947	NT	Mus musculus acetylcholine receptor beta (Acb), mRNA
4953	17679		1.02	5.2E-01	7106444	NT	Mus musculus vanilloid receptor-like protein 1 (V11), mRNA
55567	18304	31272	0.87	5.2E-01	AA284261.1	EST_HUMAN	zr44d09.77 Seires, senescent fibroblasts NbHSE Homo sapiens cDNA clone IMAGE:325169 3'
96330	25126	35474	0.75	5.2E-01	X02218.1	NT	Chicken duplicated genes for Histone H2A, H4 and a histone H3 gene
96330	25126	35476	0.75	5.2E-01	X02218.1	NT	Chicken duplicated genes for Histone H2A, H4 and a histone H3 gene
98332	22483	35685	0.48	6.2E-01	AA194818.1	EST_HUMAN	zq05bd9.r1 Stratagene muscle 837209 Homo sapiens cDNA clone IMAGE:328783 5'
99226	22574	35772	1.35	5.2E-01	AF143852.2	NT	Homo sapiens PELOTA (PELOTA) gene, complete cds
12744	25010		7	5.2E-01	P18518	SWISSPROT	RETINOIC ACID RECEPTOR GAMMA (RAR-GAMMA) (RETINOIC ACID RECEPTOR DELTA) (RAR-DELTA)
603	13381	28013	1.84	5.1E-01	M58509.1	NT	Human adrenodoxin reductase gene, exons 3 to 12
633	13412	28647	4.49	5.1E-01	AJ238944.1	NT	Polyengium Vitellinum (strain PI VI) 18S rRNA gene
633	13412	28048	4.49	5.1E-01	AJ238944.1	NT	Polyengium Vitellinum (strain PI VI) 18S rRNA gene
18448	14384		1.08	5.1E-01	X87885.1	NT	Ranvierius mRNA for mammalian fusca protein
20117	14752		1.29	5.1E-01	BF683085.1	EST_HUMAN	602139318F-1 NIH MGIC_48 Homo sapiens cDNA clone IMAGE:4288117 5'
4057	16802	229433	3.88	5.1E-01	AI858485.1	EST_HUMAN	w38p12x1 NCI OGAP-101 Homo sapiens cDNA clone IMAGE:2427283 3'
4164	16904	28533	2.81	5.1E-01	P86380	SWISSPROT	TRANSCRIPTION-REPAIR COUPLING FACTOR (TRCF)
6103	17821	30433	1.01	5.1E-01	U72863.1	NT	Human alpha 1a adrenergic receptor (alpha1a) gene, 5' flanking region
6128	18806	31874	0.97	5.1E-01	BE541068.1	EST_HUMAN	6010838608F-1 NIH MGIC_10 Homo sapiens cDNA clone IMAGE:3450000 5'
6183	18806		0.93	5.1E-01	AV712326 DGA Homo sapiens cDNA clone DCAAUFR7 5'	EST_HUMAN	AV712326 DGA Homo sapiens cDNA clone DCAAUFR7 5'
6818	19478	32502	1.69	5.1E-01	R88873.1	EST_HUMAN	yf44e09.s Seires placenta NB24P Homo sapiens cDNA clone IMAGE:146872 3'
8470	21162	34504	0.63	5.1E-01	AW806881.1	EST_HUMAN	QV4-ST0023-160400-172-e01 ST0023 Homo sapiens cDNA
8470	21162	34305	0.63	5.1E-01	AW806881.1	EST_HUMAN	QV4-ST0023-160400-172-e01 ST0023 Homo sapiens cDNA
9583	22238	35420	4.33	5.1E-01	J05412.1	NT	Human regenerating protein (reg) gene, complete cds
9587	22240	35424	3.14	5.1E-01	W222302.1	EST_HUMAN	65B1 Human retina cDNA Tap509-cleaved sublibrary Homo sapiens cDNA not direction
100060	22708	35926	0.89	5.1E-01	M94579.1	NT	Human carboxy ester lipase (CEL) gene, complete cds
12086	25137		4.28	5.1E-01	BF030207.1	EST_HUMAN	6015568635F-1 NIH MGIC_58 Homo sapiens cDNA clone IMAGE:3826767 5'
12326	24745		3.55	5.1E-01	BF439982.1	EST_HUMAN	ne65f103 NC_ CGAA_Bm23 Homo sapiens cDNA clone IMAGE:3408218 3' similar to contains element TAR1 repetitive element;
2130	14861	27590	1.24	5.0E-01	48855532	NT	Homo sapiens postmetiotic segregation Increased 2-like 9 (PMS2.9), mRNA
2130	14861	27591	1.24	5.0E-01	48855532	NT	Homo sapiens postmetiotic segregation Increased 2-like 9 (PMS2.9), mRNA

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**Table 4**  
**Single Exon Probes Expressed In Brain**

Probe Seq ID	Exon No:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
2140	14870	27801	3.19	5.0E-01	AF008210.1	NT	Buchnera aphidicola genomic fragment containing (chaperone Hsp60) groEL, DNA biosynthesis initiating protein (dsba), ATP operon (atpCDGAHFEB), and putative chromosome replication protein (gldA) genes, complete cds; and termination factor Rho (rho) gene>
2140	14870	27802	3.18	5.0E-01	AF008210.1	NT	Buchnera aphidicola genomic fragment containing (chaperone Hsp60) groEL, DNA biosynthesis initiating protein (dsba), ATP operon (atpCDGAHFEB), and putative chromosome replication protein (gldA) genes, complete cds; and termination factor Rho (rho) gene>
3811	16583	29198	1.13	5.0E-01	L38483.1	NT	Rattus norvegicus legged protein mRNA, complete cds
38864	16604	29241	2.75	5.0E-01	AB033010.1	NT	Homo sapiens mRNA for KIAA1184 protein, partial cds
6547	18312		0.86	5.0E-01	BF576189.1	EST_HUMAN	602132642F1 NIH_MGC_81 Homo sapiens cDNA clone IMAGE:4271639 5'
7562	20232	33334	0.75	5.0E-01	AL161649.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 49
7562	20232	33335	0.75	5.0E-01	AL161549.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 49
8428	21121		1.82	5.0E-01	M62304.1	NT	Xenopus laevis smooth muscle beta-tropomyosin mRNA, complete cds
8589	21261	94399	0.71	5.0E-01	BF107848.1	EST_HUMAN	601823830R1 NIH_MGC_79 Homo sapiens cDNA clone IMAGE:403485 3'
8588	20429	33647	2.74	5.0E-01	BF317212.1	EST_HUMAN	601903871F1 NIH_MGC_18 Homo sapiens cDNA clone IMAGE:4136832 5'
9525	22178	35362	1.38	6.0E-01	P35573	SWISSPROT	GLYCOCEN DEBRANCHING ENZYME (GLYCOGEN DEBRANCHER) [INCLUDES: 4-ALPHA-GLUCANO TRANSFERASE (OLIGO-1,4-1,4-GLUCANTRANSFERASE); AMYLO-1,6-GLUCOSIDASE (DEXTRIN 6-ALPHA-D-GLUCOSIDASE)]
9525	22178	35363	1.36	6.0E-01	P35573	SWISSPROT	GLYCOCEN DEBRANCHING ENZYME (GLYCOGEN DEBRANCHER) [INCLUDES: 4-ALPHA-GLUCANO TRANSFERASE (OLIGO-1,4-1,4-GLUCANTRANSFERASE); AMYLO-1,6-GLUCOSIDASE (DEXTRIN 6-ALPHA-D-GLUCOSIDASE)]
10280	22938		1.12	6.0E-01	BE868218.1	EST_HUMAN	601445024F1 NIH_MGC_85 Homo sapiens cDNA clone IMAGE:3849438 5'
12026	24654		4	5.0E-01	AF028215.1	NT	Mus musculus MRC OX-2 antigen homolog gene, exons 2-5, and complete cds
12715	24989		1.86	5.0E-01	AL1683302.2	NT	Homo sapiens chromosome 21 segment HS21C102
12726	24997		4.39	5.0E-01	O139861	SWISSPROT	NUCLEAR ENVELOPE PROTEIN CUT11
772	13544	28205	2.43	4.9E-01	BF571462.1	EST_HUMAN	602076849F1 NIH_MGC_62 Homo sapiens cDNA clone IMAGE:2243860 5'
1656	14402	27080	1.54	4.9E-01	AJ243955.1	NT	Xenopus laevis mRNA for $\alpha$ -Jun protein, 1978 BP
1899	14636	27345	1.16	4.9E-01	U40589.1	NT	Cavia porcellus pulmonary surfactant protein A (SP-a) mRNA, complete cds
5321	18124	30753	0.89	4.9E-01	Q161554	SWISSPROT	FIBRILLIN 1 PRECURSOR
5946	18728	31686	3.05	4.9E-01	AF020831.1	NT	Homo sapiens diacylglycerol kinase 3 (DAGK3) gene, exon 10
5946	18728	31687	3.05	4.9E-01	AF020831.1	NT	Homo sapiens diacylglycerol kinase 3 (DAGK3) gene, exon 10
7352	20033	33111	1.61	4.9E-01	AB04051.1	NT	Oryza sativa subsp. japonica mEF-G mRNA for mitochondrial elongation factor G, complete cds
7605	20271	33378	0.84	4.9E-01	Q10506	SWISSPROT	PUTATIVE UNDECAPRENYL-PHOSPHATE ALPHAN-ACETYLGLUCOSAMINYLTRANSFERASE
7605	20271	33379	0.84	4.9E-01	Q10506	SWISSPROT	PUTATIVE UNDECAPRENYL-PHOSPHATE ALPHAN-ACETYLGLUCOSAMINYLTRANSFERASE
8888	21579		1.45	4.9E-01	BF209791.1	EST_HUMAN	601874984F1 NIH_MGC_54 Homo sapiens cDNA clone IMAGE:4102503 5'

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Table 4  
Single Exon Probes Expressed In Brain

Probe SEQ ID NO:	Exon seq ID NO:	ORF seq ID NO:	Expression Signal	Meet Similar (T <sub>cop</sub> ) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
9086	21775	34939	0.89	4.8E-01	AW338905.1	EST_HUMAN	hs90-02.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:28072883 3' similar to TR:085714
9186	25431		1.88	4.8E-01	108468853	NT	Mus musculus unc13 homolog (C. elegans) 1 (Unc13hn), mRNA
102116	22884	36078	0.84	4.8E-01	AF053890.1	NT	Mus musculus adenylyl cyclase 1 (Adcy1) cDNA, partial cds
104119	23065	36288	0.77	4.8E-01	XSP0000.1	NT	H.sapiens DNA for BCL7A gene and BCL7A/[CH] locus fusion
119225	24486		1.72	4.8E-01	AF176912.1	NT	Homo sapiens neurotrophin-1/B-cell stimulating factor-3 gene, complete cds
12709	25392		6.73	4.8E-01	AA613632.1	EST_HUMAN	np22a11.s1 NCI CGAP Cof01 Homo sapiens cDNA clone IMAGE:1144652 3'
4288	17037		0.77	4.8E-01	4504890	NT	Homo sapiens potassium channel, subfamily K, member 5 (TASK-2) (KCNK5) mRNA, and translated products
5420	18219	30630	10.78	4.8E-01	J02887.1	NT	Saccharomyces cerevisiae) sporulation protein (SPO11) gene required for meiotic recombination, complete cds
6579	19342	32386	0.78	4.8E-01	U92882.1	NT	Mus musculus slow skeletal muscle troponin T (Tmrt1) gene, complete cds
6589	19352		3.79	4.8E-01	AA559878.1	EST_HUMAN	nsn08.s1 NCI CGAP Alv1 Homo sapiens cDNA clone IMAGE:1217513
7218	18801		1.99	4.8E-01	6031980	NT	Homo sapiens reproduction 8 (D8S2288E) mRNA
7585	20235	33339	0.78	4.8E-01	AL163208.2	NT	Homo sapiens chromosome 21 segment HS21C009
7681	20325	33424	4.05	4.8E-01	AL161492.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 4
7681	20325	33425	4.05	4.8E-01	AL161492.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 4
7805	20500	33521	1.2	4.8E-01	AI820744.1	EST_HUMAN	Y7710.5 Soares breast 2NbbH81 Homo sapiens cDNA clone IMAGE:154785 5' similar to contains element MER8 repetitive element;
8144	21875		0.92	4.8E-01	BE155148.1	EST_HUMAN	PM1+HT0350-201289-004-504 HT0350 Homo sapiens cDNA S.cerevisiae ORFs from chromosome X
10529	23322		1.88	4.8E-01	X83502.1	NT	Typanosoma cruzi transposon VIP II/SIRE repeat region
12217	25165		3.04	4.8E-01	AF227565.1	NT	Chlamydomonas reinhardtii cop gene, exons 1-3
12786	26216		1.88	4.8E-01	AJ132684.1	NT	Chlamydomonas reinhardtii cop gene, exons 1-3
6422	19190	32188	8.41	4.7E-01	BF217173.1	EST_HUMAN	6018836801F1 NIH_MCG_57 Homo sapiens cDNA clone IMAGE:40963387 5'
6841	19423	32438	0.84	4.7E-01	AI204374.1	EST_HUMAN	ct72a08.x1 Soares testis_NIH Homo sapiens cDNA clone hbc811 5' end
7784	20460	33684	0.83	4.7E-01	T11414.1	EST_HUMAN	hbc811 Human pancreatic islet Homo sapiens cDNA clone hbc811 5' end
7784	20480	33585	0.83	4.7E-01	T11414.1	EST_HUMAN	hbc811 Human pancreatic islet Homo sapiens cDNA clone hbc811 5' end
8874	21684	34816	0.52	4.7E-01	6981501	NT	Rattus norvegicus Spermine binding protein (Sbp), mRNA
10751	23428		6.11	4.7E-01	AF102873.1	NT	Influenza A virus isolate Hk51697 hemagglutinin (H7) gene, partial cds
11022	23894	36957	2.2	4.7E-01	U41058.1	NT	Human collagen alpha2(XI) (COL11A2) gene, exons 8 through 16, and partial cds
11252	23914	37208	1.61	4.7E-01	BF528858.1	EST_HUMAN	6002043889FI NCI CGAP Brm87 Homo sapiens cDNA clone IMAGE:4181303 5'
11349	24039	37342	1.7	4.7E-01	AW889448.1	EST_HUMAN	RC8-NT0029-240400-011-E08 NT0029 Homo sapiens cDNA
12116	24609		1.52	4.7E-01	BE887763.1	EST_HUMAN	601611333FI NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3912488 5'
12237	24689		1.51	4.7E-01	AW341561.1	EST_HUMAN	Ind11c08.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2809188 3'

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## Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
12813	25055			1.63	4.7E-01 AF000007.1	NT	<i>Pyrococcus horikoshii</i> OT3 genomic DNA, 1495001-1738505 nt, position 777
12817	25300			1.38	4.7E-01 0879502	NT	Mus musculus proteasome (prosome, macrophain) 26S subunit, ATPase 3 (Psmc3), mRNA
12823	16479	29116		1.57	4.6E-01 BF589300.1	EST_HUMAN	<i>Bacillus</i> 143F1 NIH MGC_81 Homo sapiens cDNA clone IMAGE:4245481 5'
3726	16479	29117		1.57	4.6E-01 BF589300.1	EST_HUMAN	<i>Bacillus</i> 143F1 NIH MGC_81 Homo sapiens cDNA clone IMAGE:4245481 5'
5333	18136	30795		1	4.6E-01 BF513593.1	EST_HUMAN	<i>Bacillus</i> 193F1 NIH MGC_19 Homo sapiens cDNA clone IMAGE:4128472 5'
5333	18138	30798		1	4.6E-01 BF513593.1	EST_HUMAN	<i>Bacillus</i> 193F1 NIH MGC_19 Homo sapiens cDNA clone IMAGE:4128472 5'
5385	18185	30875		3.11	4.6E-01 Q90849	SWISSPROT	INTERFERON REGULATORY FACTOR 3 (IRF-3)
5385	18185	30876		3.11	4.6E-01 Q90849	SWISSPROT	INTERFERON REGULATORY FACTOR 3 (IRF-3)
6459	18258	31148		1.84	4.6E-01 BE734781.1	EST_HUMAN	801568755F1 NIH MGC_21 Homo sapiens cDNA clone IMAGE:3843837 5'
5472	18271	31163		2.17	4.6E-01 AI247079.1	EST_HUMAN	q5f50j02.x1 Soares, fetal liver_spleen, INF/S_S1 Homo sapiens cDNA clone IMAGE:18490113 similar to TR-Q15338 O15338 BUTYROPHILIN.
5472	18271	31164		2.17	4.6E-01 AI247079.1	EST_HUMAN	q5f50j02.x1 Soares, fetal liver_spleen, INF/S_S1 Homo sapiens cDNA clone IMAGE:18490113 similar to TR-Q15338 BUTYROPHILIN.
6480	18278	31175		1.6	4.6E-01 P20050	SWISSPROT	MEIOSIS SPECIFIC PROTEIN HOP1
5560	18357			0.98	4.6E-01 AF2412124.1	NT	<i>Andrea schwartzii</i> Cytocrome b gene, partial cds; mitochondrial gene for mitochondrial product
5645	18440			0.77	4.6E-01 BE817247.1	EST_HUMAN	PM0-BN0260-120300-001-F07 BN0260 Homo sapiens cDNA
6809	18598	31528		0.59	4.6E-01 D26215.1	NT	Unidentified soil bacteria 16S rRNA gene encoding 16S ribosomal RNA
6163	18540	31911		1.21	4.6E-01 AE000894.1	NT	<i>Methanobacterium thermophilicum</i> from bases 1165751 to 1176238 (section 100 of 148) of the complete genome
6669	19586	32620		3.2	4.6E-01 U62232.1	NT	<i>Emericella nidulans</i> NEMPA (nemPA) gene, mitochondrial gene encoding putative mitochondrial protein, complete cds
6669	19588	32621		3.2	4.6E-01 U62232.1	NT	<i>Emericella nidulans</i> NEMPA (nemPA) gene, mitochondrial gene encoding putative mitochondrial protein, complete cds
7131	25105	32884		0.57	4.6E-01 L07320.1	NT	<i>Marine cyanobacteria</i> s1 protein gene, complete cds
7629	20285	33403		0.91	4.6E-01 AA493577.1	EST_HUMAN	rh0405.s1 NCI CG3AP_Thy1 Homo sapiens cDNA clone IMAGE:943353 similar to contains Alu repetitive elementcontains element L1 repetitive element;
7658	20322			0.59	4.6E-01 Q80069	SWISSPROT	GENOME POLYPROTEIN [CONTAINS: N-TERMINAL PROTEIN (P1); HELPER COMPONENT PROTEINASE (HC-PRO); PROTEIN P3; 6 KD PROTEIN 1 (6K1); CYTOPLASMIC INCLUSION PROTEIN (C1); 8 KD PROTEIN 2 (8K2); GENOME-LINKED PROTEIN (VPG); NUCLEAR INCLUSION PROTEIN A (NI-A) (N1-A)]
8219	20913	34049		10.11	4.6E-01 BF89739.1	EST_HUMAN	602130935F1 NIH MGC_58 Homo sapiens cDNA clone IMAGE:4227828 5'
9201	21870	35035		1.11	4.6E-01 P55202	SWISSPROT	ATRIAL NATRIURETIC PEPTIDE RECEPTOR B PRECURSOR (ANP-B) (ANPRB) (GC-B) (GUANYLATE CYCLASE)

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**Table 4**  
**Single Exon Probes Expressed In Brain**

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source.	Top Hit Descriptor
9201	21870	35038	1.11	4.0E-01	P55202	SWISSPROT	ATRIAL NATRIURETIC PEPTIDE RECEPTOR B PRECURSOR (ANP-B) (ANPRB) (GC-B) (GUANYLATE CYCLASE)
9878	22528	35720	1.64	4.0E-01	AI915634.1	EST_HUMAN	wg73e12.x1 Soares_NSF_F8_8W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:2370768 3'
9878	22528	35721	1.64	4.0E-01	AI915634.1	EST_HUMAN	wg73e12.x1 Soares_NSF_F8_8W_OT_PA_P_S1 Homo sapiens cDNA clone IMAGE:2370768 3'
10912	23592		2.3	4.0E-01	P88163	SWISSPROT	PUTATIVE VITELLOGENIN RECEPTOR PRECURSOR (YL)
10922	23602	36850	10.22	4.0E-01	BE185449.1	EST_HUMAN	IL5-HT0730-100500-075-005 HT0730 Homo sapiens cDNA
10922	23602	36851	10.22	4.0E-01	BE185449.1	EST_HUMAN	IL6-HT0730-100500-076-006 HT0730 Homo sapiens cDNA
11450	23217	36449	5.32	4.0E-01	AF01889.1	NT	Human thiopurine methyltransferase (TPMT) gene, exon 10 and complete cds
11450	23217	36450	5.32	4.0E-01	AF01889.1	NT	Human thiopurine methyltransferase (TPMT) gene, exon 10 and complete cds
12163	24845		1.77	4.0E-01	D53316.1	EST_HUMAN	HUM105FF03B Clontech human fetal brain polyA+ mRNA (#86355) Homo sapiens cDNA clone GEN-105FF03
13004	14641	27350	1.43	4.0E-01	AE001831.1	NT	Dermococcus radiodurans R1 section 68 of the complete chromosome 1
13004	14641	27351	1.43	4.0E-01	AE001831.1	NT	Dermococcus radiodurans R1 section 68 of the complete chromosome 1
28773	15840	28284	4.5	4.0E-01	AA677086.1	EST_HUMAN	2165d02.s1 Soares_fetal_liver_spleen_1NFLS_S1 Homo sapiens cDNA clone IMAGE:454179 3'
33112	18072	28722	4.58	4.0E-01	Q05783	SWISSPROT	BASEMENT MEMBRANE-SPECIFIC HEPARAN SULFATE PROTEOGLYCAN CORE PROTEIN PRECURSOR (HSPG) (PERLECAN) (PLC)
3372	18191	28787	1.07	4.0E-01	AF128378.1	NT	Mus musculus DNA polymerase epsilon catalytic subunit (Pole) gene, exons 2 through 12
4007	16753		0.95	4.0E-01	Q28247	SWISSPROT	COLLAGEN ALPHA 5(IV) CHAIN
40555	16890	28431	0.88	4.0E-01	AI708908.1	EST_HUMAN	ss86e08.x1 Barshardt et al. HPLB6 Homo sapiens cDNA clone IMAGE:2353480 3'
41155	17287		4.25	4.0E-01	AW873495.1	EST_HUMAN	hg80302.x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:3041810 3'
4890	17617	30236	1.1	4.0E-01	BE862445.2	EST_HUMAN	601657225RT NIH_MGC_67 Homo sapiens cDNA clone IMAGE:3886023 3'
54481	18280	31161	1.49	4.0E-01	AW609814.1	EST_HUMAN	QV2zP10012-140100-031-008 PT0012 Homo sapiens cDNA
6510	19276		1.46	4.0E-01	Q00956	SWISSPROT	COAT PROTEIN
7312	18995	33073	1.27	4.0E-01	M37036.1	NT	Pat nucleolar proteins B23.1 and B23.2
7609	20180	38273	2.64	4.0E-01	AI858849.1	EST_HUMAN	wf32a02.x1 NCI_OGAP_U1 Homo sapiens cDNA clone IMAGE:2426818 3' similar to TR-Q928223 Q92923
7821	20287	33398	0.85	4.0E-01	P50070	SWISSPROT	SWI/SNF COMPLEX 170 KDA SUBUNIT
8206	20800		0.88	4.0E-01	M32981.1	NT	D.melanogaster Shm2 protein mRNA, complete cds
8302	20896	34134	3.5	4.0E-01	AI64886.1	EST_HUMAN	tz6fg11.x1 NCI_CGAP_Ov35 Homo sapiens cDNA clone IMAGE:2202844 3'
8457	21149	34292	0.83	4.0E-01	Q52728	SWISSPROT	POLY-BETA-HYDROXYBUTYRATE POLYMERASE (POLY(3-HYDROXYBUTYRATE) POLYMERASE) (PHB POLYMERASE) (PHB SYNTHASE) (POLY(3-HYDROXYALKANOATE) POLYMERASE) (PHA SYNTHASE) (POLYHYDROXYALKANOIC ACID SYNTHASE)
8680	21372		2.34	4.0E-01	11444786.1	NT	Homo sapiens hypothetical protein DK7zP547G183 (DK7zP547G183) mRNA

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
8897	21588	34728	0.88	4.5E-01	AE000218.1	NT	Escherichia coli K-12 MG1655 section 108 of the complete genome
8840	22491		1.02	4.5E-01	96309816	NT	Bombyx mori nuclear polyhedrosis virus, complete genome
10392	23038	36254	24.62	4.5E-01	M88003.1	EST_HUMAN	EST02531 Fetal brain, Strategene (cat#936206) Homo sapiens cDNA clone HFBCY17
10392	23038	36255	24.62	4.5E-01	M88003.1	EST_HUMAN	EST02531 Fetal brain, Strategene (cat#936206) Homo sapiens cDNA clone HFBCY17
10772	23455	36898	2.15	4.5E-01	AW891271.1	EST_HUMAN	x014h01_x1_NCL_CGAP_U13 Homo sapiens cDNA clone IMAGE:2703985 3' similar to SW:INT6_MOUSE Q84252 VIRAL_INTEGRATION SITE PROTEIN INT-6 [1];
11217	23850		1.52	4.5E-01	AV718382.1	EST_HUMAN	AV718382 GLC Homo sapiens cDNA clone GLCCED12 5'
11895	25384		3.52	4.5E-01	BE871481.1	EST_HUMAN	601449201_F1_NIH_MGC_65 Homo sapiens cDNA clone IMAGE:3852981 5'
12840	24880		1.58	4.5E-01	BF337831.1	EST_HUMAN	60203275_F1_NCL_CGAP_Bm84 Homo sapiens cDNA clone IMAGE:4183280 5'
12811	24918		3.37	4.5E-01	11422089	NT	Homo sapiens testis-specific kinase 2 (TESK2), mRNA
2388	15109	27847	3.39	4.4E-01	P49785	SWISSPROT	VASCULAR ENDOTHELIAL GROWTH FACTOR B PRECURSOR (VEGF-B) (VEGF RELATED FACTOR)
3510	16070	28719	1.29	4.4E-01	AF058790.1	NT	Rattus norvegicus SynGAP-6 mRNA, complete cds
3510	16070	28720	1.29	4.4E-01	AF058790.1	NT	Rattus norvegicus SynGAP-6 mRNA, complete cds
3513	16073	28723	2.92	4.4E-01	BF056728.1	EST_HUMAN	797d021_f1_NCL_CGAP_B118 Homo sapiens cDNA clone IMAGE:3393795 5'
42039	16550		1.88	4.4E-01	BE378707.1	EST_HUMAN	60123713_F1_NIH_MGC_44 Homo sapiens cDNA clone IMAGE:3603693 5'
63334	18137	30767	1.2	4.4E-01	P04829	SWISSPROT	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR
63334	18137	30793	1.2	4.4E-01	P04829	SWISSPROT	HISTIDINE-RICH GLYCOPROTEIN PRECURSOR
56022	18397	31309	1.59	4.4E-01	S85019.1	NT	muclin [rats, Sprague-Dawley, sulfur-dioxide-treated tracheal epithelium, mRNA Partial 380 nt]
65119	18415	31323	2	4.4E-01	AV720408.1	EST_HUMAN	AV720408 GLC Homo sapiens cDNA clone GLCCSC12 5'
6884	18851	31591	1.46	4.4E-01	AI188413.1	EST_HUMAN	qf62h11_x1_NCL_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:1861125 3' similar to TR:Q28168 Q28168 UNKNOWN PROTEIN;
6884	18851	31592	1.48	4.4E-01	AI188413.1	EST_HUMAN	qf62h11_x1_NCL_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:1861125 3' similar to TR:Q28168 Q28168 UNKNOWN PROTEIN;
6146	18923	31894	1.78	4.4E-01	AW080795.1	EST_HUMAN	x271e08_x1_NCL_CGAP_Co18 Homo sapiens cDNA clone IMAGE:2585510 3' similar to TR:Q88154 Q88154 AFLATOXIN B1-ALDEHYDE REDUCTASE ;
6236	18010		1.42	4.4E-01	AA778132.1	EST_HUMAN	aa86d11_s1_Stratagene seattle brain S11 Homo sapiens cDNA clone IMAGE:970985 3' similar to gb:M16038 TYROSINE-PROTEIN KINASE LYN (HUMAN);
7287	18880	33058	1.04	4.4E-01	AE000571.1	NT	Helicobacter pylori 26895 section 48 of 134 of the complete genome
7723	25119		0.8	4.4E-01	AE001198.1	NT	Tsponeroma pallidum section 4 of 87 of the complete genome
7740	20438		9.71	4.4E-01	Z11678.1	NT	S.tuberorum mRNA for induced stolon tip protein (partial)
8681	21353	34500	0.84	4.4E-01	AA056427.1	EST_HUMAN	Z68903_s1_Stratagene clone (#837204) Homo sapiens cDNA clone IMAGE:5098386 3'
8049	21738	34896	0.7	4.4E-01	AF412640.1	NT	HIV-1 isolates 081076 from USA, envelope glycoprotein (env) gene, partial cds

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
8082	21771	34834	0.57	4.4E-01	AW612578.1	EST_HUMAN	h195cg8.x1 NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE:2854222 3' similar to SW:MSH6_HUMAN_P52701 DNA MISMATCH REPAIR PROTEIN MSH6 :
9180	21860	35025	1.24	4.4E-01	062838	SWISSPROT	ZINC FINGER X-CHROMOSOMAL PROTEIN
8862	22512	35709	2.18	4.4E-01	AI288850.1	EST_HUMAN	qg39f09.x1 NCI_CGAP_Lif5 Homo sapiens cDNA clone IMAGE:1810821 3'
8863	22513		1.98	4.4E-01	P28922	SWISSPROT	GLYCOPROTEIN B PRECURSOR (GLYCOPROTEIN 14)
8867	22845	35857	4.31	4.4E-01	P35590	SWISSPROT	TYROSINE-PROTEIN KINASE RECEPTOR TIE-1 PRECURSOR
10273	22821	36132	1.33	4.4E-01	S78404.1	NT	Beta-HKA-H,K-ATPase beta-subunit [rat], Genomic, 6883 nt, segment 2 of 2
10273	22821	36133	1.33	4.4E-01	S78404.1	NT	Beta-HKA-H,K-ATPase beta-subunit [rat], Genomic, 6883 nt, segment 2 of 2
12448	246825	31095	3.44	4.4E-01	6877874	NT	Mus musculus sodium channel, type X, alpha polypeptide [Scn1Ba], mRNA
12679	246903	31000	3.36	4.4E-01	9627742	NT	Autographa californica nucleopolyhedrovirus, complete genome
12883	24971		1.91	4.4E-01	P54725	SWISSPROT	UV EXCISION REPAIR PROTEIN RAD23 HOMOLOG A (HHR23A)
12786	25152		1.43	4.4E-01	AW363338.1	EST_HUMAN	RC2-CT0320-281169-012-007 CT0320 Homo sapiens cDNA
402	13187	25835	2.17	4.3E-01	AF158218.1	NT	Callithrix jacchus MW/LW opsin gene, upstream flanking region
402	13187	25836	2.17	4.3E-01	AF158218.1	NT	Callithrix jacchus MW/LW opsin gene, upstream flanking region
2875	15842		1.64	4.3E-01	AW935269.1	EST_HUMAN	CM2-DT0003-01020-077-001 DT0003 Homo sapiens cDNA
3058	15822	28468	0.75	4.3E-01	AW989477.1	EST_HUMAN	MFO-BN0070-270300-008-504 BN0070 Homo sapiens cDNA
4131	16873	28501	1.29	4.3E-01	J00306.1	NT	Homo somatic statin 1 gene and flanks
4374	13187	25835	1.18	4.3E-01	AF158218.1	NT	Callithrix jacchus MW/LW opsin gene, upstream flanking region
4374	13187	25836	1.18	4.3E-01	AF158218.1	NT	Callithrix jacchus MW/LW opsin gene, upstream flanking region
4902	17629		1.19	4.3E-01	AL161502.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 14
5280	18085	30742	0.33	4.3E-01	P48634	SWISSPROT	LARGE PROLINE-RICH PROTEIN BA12 (HLA-B-ASSOCIATED TRANSCRIPT 2)
5280	18085	30743	0.3	4.3E-01	P48634	SWISSPROT	LARGE PROLINE-RICH PROTEIN BAT2 (HLA-B-ASSOCIATED TRANSCRIPT 2)
5793	18589	91515	1.69	4.3E-01	BE181855.1	EST_HUMAN	QV1-HT0639-070500-191-d08 HT0638 Homo sapiens cDNA
6817	18609	31634	2.02	4.3E-01	AF78825.1	NT	Selemtid selemus olfactory receptor (SSC180) gene, partial cds
6838	19371	32284	4.73	4.3E-01	AJ001678.1	NT	Catolymn cotonicus leponica IfnQ gene
6839	19606	322848	0.3	4.3E-01	AF075628.1	NT	Equus caballus microsatellite LEX027
6707	19511		0.91	4.3E-01	O38367	SWISSPROT	DNA GYrase SUBUNIT B
7329	20011		1.88	4.3E-01	BF348001.1	EST_HUMAN	602023134F1 NCI_CGAP_Bm67 Homo sapiens cDNA clone IMAGE:4168283 5'
7496	20168	33280	0.61	4.3E-01	U51002.1	NT	Mus musculus Dik-2 gene, complete cds
8326	21019		2.72	4.3E-01	J87040.1	NT	Methanococcus voltae flagella-related protein C-1 (flaC-f1a) genes, complete cds
9154	21885	35053	0.98	4.3E-01	Y14604.1	NT	Ermifia amylovora resV gene
9826	22279	35468	2.18	4.3E-01	AW630048.1	EST_HUMAN	Itt74e10.y1 NCI_CGAP_G11 Homo sapiens cDNA clone IMAGE:2868854 5'
8626	22279	35469	2.18	4.3E-01	AW630048.1	EST_HUMAN	Itt74e10.y1 NCI_CGAP_G11 Homo sapiens cDNA clone IMAGE:2868854 5'

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Single Exon Probes Expressed in Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Meet Similar (Top) Hit BLAST E Value	Top Hit No.	Top Hit Database Source	Top Hit Descriptor
10128	22776	35890	0.84	4.3E-01 AW170559.1	EST_HUMAN	xt63e05.x1 Soares_NHCeC cervical_tumor Homo sapiens cDNA clone IMAGE:26998400 3' similar to TRC001189_000189_MU-ADAPTIN-RELATED PROTEIN 2;	
10409	23055	36272	0.5	4.3E-01 H65282.1	EST_HUMAN	y45b05.s1 Soares_fetal liver spleen INFSL Homo sapiens cDNA clone IMAGE:208209 3'	
10849	19608	32646	2.45	4.3E-01 AF075528.1	NT	Equus caballus microsatellite LEX027	
11168	23833	37113	1.29	4.3E-01 AW882658.1	EST_HUMAN	RC3-BN0034-280200-013-c12 BN0034 Homo sapiens cDNA	
11168	23833	37114	1.29	4.3E-01 AW882658.1	EST_HUMAN	RC3-BN0034-280200-013-c12 BN0034 Homo sapiens cDNA	
11745	24336	37862	1.84	4.3E-01 AI874322.1	EST_HUMAN	h84d04.x1 NCI_CGAP_Ox35 Homo sapiens cDNA clone IMAGE:2283351 3'	
12770	25025		2.18	4.3E-01 AJ003022.1	NT	Streptomyces coelicolor wrH gene	
1337	15568	26761	1.64	4.2E-01 Q39102	SWISSPROT	CELL DIVISION PROTEIN FISH HOMOLOG PRECURSOR	
1941	14878		1.23	4.2E-01 AA761653.1	EST_HUMAN	n24a08.s1 NCI_CGAP_GCB11 Homo sapiens cDNA clone IMAGE:1288988 3'	
3599	16349	28890	4.4	4.2E-01 AE003947.1	NT	Xylella fastidiosa, section 83 of 228 of the complete genome	
3628	16381	28021	1.41	4.2E-01 AI280338.1	EST_HUMAN	q194b01.x1 Soares_NhIMpu_S1 Homo sapiens cDNA clone IMAGE:1878945 3'	
3698	17898		0.85	4.2E-01 N81203.1	EST_HUMAN	7381ET fetal brain cDNA Homo sapiens cDNA clone 7881ET-K similar to R07879_Z40498	
3964	18713	28352	0.97	4.2E-01 Q04888	SWISSPROT	SOX-8 PROTEIN	
4649	17383	30016	4.88	4.2E-01 AA534093.1	EST_HUMAN	hj69h01.s1 NCI_CGAP_PR10 Homo sapiens cDNA clone IMAGE:997777 similar to gbaM333600 HLA CLASS II HISTOCOMPATIBILITY ANTIGEN, DR-1 BETA CHAIN (HUMAN);	
4731	17463	30100	3.48	4.2E-01 R13487.1	EST_HUMAN	yf77e01.r1 Soares_infant brain 1NIB Homo sapiens cDNA clone IMAGE:28278 5'	
5626	18423	31338	0.82	4.2E-01 BF242055.1	EST_HUMAN	60187872/11FT NIH_MGC_55 Homo sapiens cDNA clone IMAGE:4108483 5'	
5693	18487	31408	1.53	4.2E-01 AW854162.1	EST_HUMAN	RC3-CT0284-060400-029-904 CT0284 Homo sapiens cDNA	
6112	18889	31858	1.01	4.2E-01 AL183247.2	NT	Homo sapiens chromosome 21 segment HS21C047	
6552	19552	32582	10.8	4.2E-01 AU158472.1	EST_HUMAN	AU188472 PLACE22 Homo sapiens cDNA clone PLACE2000470 3'	
6552	19552	32583	10.8	4.2E-01 AU158472.1	EST_HUMAN	AU158472 PLACE22 Homo sapiens cDNA clone PLACE2000470 3'	
6911	25101	32894	2.16	4.2E-01 S82504.1	NT	Bircan=breast cancer gene [rats, WF, spleen, Gerarts, 419 nt, segment 2 of 2]	
6933	19886	32734	7	4.2E-01 AL181547.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 47	
7891	20586	33715	2.21	4.2E-01 AW867448.1	EST_HUMAN	ES1368413 MAGE resequences, MAGE Homo sapiens cDNA	
7891	20586	33716	2.21	4.2E-01 AW857448.1	EST_HUMAN	ES1368413 MAGE resequences, MAGE Homo sapiens cDNA	
8108	20800	33932	0.61	4.2E-01 4758039	NT	Homo sapiens cytochrome c oxidase subunit Vic (COX6C), nuclear gene encoding mitochondrial protein, mRNA	
9870	22520		0.94	4.2E-01 AA705007.1	EST_HUMAN	z95f01.s1 Soares_fetal_liver_spleen INFSL Homo sapiens cDNA clone IMAGE:462849 3'	
1081	22729	35944	0.45	4.2E-01 AF181854.1	NT	Lassa virus strain 803213 glycoprotein precursor and nucleoprotein genes, complete cds	
10383	23039	36268	1.78	4.2E-01 AW863668.1	EST_HUMAN	MF3-SN0010-280300-103-h07 SN0010 Homo sapiens cDNA	
10972	23648	36901	2.69	4.2E-01 AB023489.1	NT	Oryzias latipes OGC7 mRNA for membrane guanylyl cyclase, complete cds	
11370	23977	37277	2.11	4.2E-01 BE864852	EST_HUMAN	601680332R1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3906085 3'	
1072	13830	26498	1.83	4.1E-01 AI905481.1	EST_HUMAN	RC-BT091-210199-142 BT091 Homo sapiens cDNA	

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Table 4  
Single Exon Probes Expressed in Brain

Probe SEQ ID NO.	Exon SEQ ID NO.	ORF SEQ ID NO:	Expression Signal	Meet Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
1081	128339	28497	1.1	4.1E-01	AV705243.1	EST_HUMAN	AV705243 ADB Homo sapiens cDNA clone ADRA-HF08 5'
1081	138339	28498	1.1	4.1E-01	AV705243.1	EST_HUMAN	AV705243 ADB Homo sapiens cDNA clone ADRA-HF08 5'
2715	15422	28181	1.1	4.1E-01	7705283 NT	Homo sapiens amphioxus-promoting complex subunit 7 (APC7), mRNA	
2841	15708	28355	2.17	4.1E-01	AL161536.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 36
2841	15708	28356	2.17	4.1E-01	AL161536.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 36
3754	16506	28142	0.68	4.1E-01	AW861292.1	EST_HUMAN	EST73384 IMAGE sequences, MAGG Homo sapiens cDNA
3754	16506	28143	0.68	4.1E-01	AW861292.1	EST_HUMAN	EST73384 IMAGE sequences, MAGG Homo sapiens cDNA
4241	16982	28607	2.93	4.1E-01	AJ249207.1	NT	Rhodococcus sp. AD45 isoC, isoH, isoJ, isoL, isoC, isoD, isoE and isoF genes
4271	17011	0.82	4.1E-01	AA809257.1	EST_HUMAN	am33-d02.51 Scores_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1542819 3'	
4618	17363	28988	1.48	4.1E-01	AV747880 NPC Homo sapiens cDNA clone NP_CDDF10 5'		
4888	18057	28706	2.48	4.1E-01	AA800344.1	EST_HUMAN	qj94b08.51 Scores_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:1505943 3'
6889	18684	31632	4.72	4.1E-01	BF681393.1	EST_HUMAN	6002166590F1 NIH_MGC_G3 Homo sapiens cDNA clone IMAGE:4297319 5'
7332	22014	33082	2.76	4.1E-01	U67686.1	NT	Methanococcus jannaschii section 77 of 150 of the complete genome
7835	22680	33177	1.38	4.1E-01	BF574804.1	EST_HUMAN	6021332826F1 NIH_MGC_S1 Homo sapiens cDNA clone IMAGE:4298238 5'
8898	21678	34827	1.39	4.1E-01	8755621 NT	Mus musculus signaling intermediate in Toll pathway-evolutionarily conserved (Sipeo-pending), mRNA	
9465	22076		0.67	4.1E-01	AF160597.1	NT	Vesicle gymnoeaudus Vgym560 cytochrome b (cytb) gene, complete cds; mitochondrial gene for mitochondrial product
10163	22811		1.05	4.1E-01	AL139078.2	NT	Campylobacter jejuni NCTC11188 complete genome; segment 3/8
10310	22857	36173	0.91	4.1E-01	AV849578.1	EST_HUMAN	AV849578 GLC Homo sapiens cDNA clone GLCBVD12 3'
10404	23050	36267	0.61	4.1E-01	P18584	SWISSPROT	PROBABLE SERINE PROTEASE DO-LIKE PRECURSOR (58 kDa IMMUNOGENIC PROTEIN) (SK69)
10404	23050	36268	0.61	4.1E-01	P18584	SWISSPROT	PROBABLE SERINE PROTEASE DO-LIKE PRECURSOR (58 kDa IMMUNOGENIC PROTEIN) (SK69)
10478	23124		1.33	4.1E-01	BF348832.1	EST_HUMAN	CN24HT0137-200888-010-008 HT0137 Homo sapiens cDNA
10743	23430	36873	80.48	4.1E-01	X58700.1	NT	Zea mays ZPMS2 gene for 19 kDa zinc protein
11368	23177	36404	2	4.1E-01	Q09470	SWISSPROT	VOLTAGE-GATED POTASSIUM CHANNEL PROTEIN KV1.1 (HUK1) (HBRK1)
12476	25360		2.62	4.1E-01	D87676.1	NT	Homo sapiens DNA for amyloid precursor protein, complete cds
1016	13775	26435	0.82	4.0E-01	8404658 NT	Lacistema rubellus mitochondrial, complete genome	
1316	14085	26739	0.95	4.0E-01	AF203478.1	NT	Drosophila melanogaster Daimantina (dm) mRNA, complete cds
1468	14216		4.05	4.0E-01	6879258 NT	Mus musculus platelet-derived growth factor receptor, beta polypeptide (Pdgfrb), mRNA	
1889	15583	27467	1.16	4.0E-01	Z86838.1	NT	Ascochyta immersus mas22 gene
1889	15583	27458	1.16	4.0E-01	Z86838.1	NT	Ascochyta immersus mas22 gene
2168	14888	27619	1.19	4.0E-01	AE011831.1	NT	Deltaproteobacter radiodurans R1 section 68 of 228 of the complete chromosome 1
2168	14888	27620	1.19	4.0E-01	AE011831.1	NT	Deltaproteobacter radiodurans R1 section 68 of 228 of the complete chromosome 1
2808	12952	25955	1.4	4.0E-01	6878430 NT	Mus musculus ubiquitin-protein ligase es3 component n-recognin (Ubr1), mRNA	

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**Table 4**  
**Single Exon Probes Expressed In Brain**

Probe SEQ ID NO:	Exon seq ID NO:	ORF SEQ ID NO:	Expression Signal	Meet Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
2868	15734	28383	1.1	4.0E-01	AL1632B0.2	NT	Homo sapiens chromosome 21 segment HS21C30
2868	15734	28384	1.1	4.0E-01	AL1632B0.2	NT	Homo sapiens chromosome 21 segment HS21C30
3683	18436	28080	1.88	4.0E-01	AF068908.1	NT	Streptococcus pneumoniae YIC (YfC), YID (YfD), penicillin-binding protein 2x (pbp2x), and undecaprenyl-phosphate-UDP-MurNAc-pentapeptide phospho-MurNAc-penta-peptide transferase (mraY) genes, complete cds
3807	16559	29191	3.38	4.0E-01	AJ277511.1	NT	Ovis aries partial JD2 gene for T cell receptor delta chain (TCRD2), exon 1
3807	16559	28192	3.38	4.0E-01	AJ277511.1	NT	Ovis aries partial JD2 gene for T cell receptor delta chain (TCRD2), exon 1
4767	17499		7.97	4.0E-01	Q31849	SWISSPROT	NADH-PLASTOQUINONE OXIDOREDUCTASE CHAIN 5, CHLOROPLAST
5820	18609	31538	1.23	4.0E-01	AW970010.1	EST_HUMAN	EST382891 MAGE sequences, MAGK Homo sapiens cDNA
6345	19116	32104	0.94	4.0E-01	P27285	SWISSPROT	STRUCTURAL POLYPROTEIN (P130) [CONTAINS: COAT PROTEIN C; SPIKE GLYCOPROTEINS E3, E2 AND E1; 6 KD PEPTIDE]
7728	20391	33504	0.88	4.0E-01	P27546	SWISSPROT	MICROTUBULE-ASSOCIATED PROTEIN 4
7829	20524	33849	0.44	4.0E-01	BF092634.1	EST_HUMAN	MR4-TN0110-160800-202-gf02 TN0110 Homo sapiens cDNA
7910	20605	33736	1.04	4.0E-01	AB016826.1	NT	Homo sapiens OCTN2 gene, complete cds
8904	21595	34736	1.17	4.0E-01	AJ323239.1	EST_HUMAN	EST26068 Cerebellum II Homo sapiens cDNA 5' end similar to EST containing Alu repeat
11680	24159		2.03	4.0E-01	BF030262.1	EST_HUMAN	601558283F1 NIH MGCG 58 Homo sapiens cDNA clone IMAGE-3828092 5'
11721	24316		2.89	4.0E-01	L76380.1	NT	Synaptosomal sp. PCC 9413 transposase gene, complete cds
12162	25222		2.26	4.0E-01	AL163302.2	NT	Homo sapiens chromosome 21 segment HS21C100
12684	24972		2.2	4.0E-01	P26049	SWISSPROT	HYPOTHETICAL 49.7 KD PROTEIN IN GIN2-ST13 INTERGENIC REGION
1356	14104	26780	1.85	3.9E-01	AF208618.1	NT	Gorilla gorilla carboxy-ester lipase (CEL) gene, complete cds
2643	15353	28101	3.34	3.9E-01	AB033019.1	NT	Homo sapiens mRNA for KIAA1183 protein, partial cds
2709	15416	28163	4.27	3.9E-01	X82032.1	NT	H. sapiens B-myb gene
2709	15416	28164	4.27	3.9E-01	X82032.1	NT	H. sapiens B-myb gene
3033	15858	28499	4.73	3.9E-01	AJ225896.1	NT	Streptozotobium mellifaciogli, syrB2, crys2 genes and of3
4039	16804	29435	1.05	3.9E-01	BF592611.1	EST_HUMAN	761401-X1 NCI_CGAP_B178 Homo sapiens cDNA clone IMAGE-3339169 3'
4832	17680	30270	1.74	3.9E-01	BE728687.1	EST_HUMAN	801583949F1 NIH MGCG_20 Homo sapiens cDNA clone IMAGE-3833669 6'
6843	18681	31568	3.81	3.9E-01	BF208036.1	EST_HUMAN	8018823262F1 NIH MGCG_53 Homo sapiens cDNA clone IMAGE-4082055 5'
7654	20549	33674	0.82	3.9E-01	U79415.1	NT	Homo sapiens prepro dipeptidyl peptidase (DPP-4) gene, complete cds
8760	21452	34800	0.81	3.9E-01	AW77011.1	EST_HUMAN	CM3-CT0105-70800-004-008 CT0105 Homo sapiens cDNA
8769	21461		0.58	3.9E-01	BF3486834.1	EST_HUMAN	802015844F1 NCI CGAP_Bm87 Homo sapiens cDNA clone IMAGE-4155322 5'
9134	21822	34988	1.26	3.9E-01	AW195898.1	EST_HUMAN	XN8604_X1 Scores_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE-2701351 3' similar to TR-093821 OBA821 KIAA0713 PROTEIN;
9445	22122	35301	1.40	3.9E-01	AI937337.1	EST_HUMAN	WP766022X1 NCI CGAP_Bm25 Homo sapiens cDNA clone IMAGE-2467658 3' similar to SW:RPX5_HUMAN P48382 BINDING REGULATORY FACTOR;

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Table 4  
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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
9778	22429	36636	3.03	3.8E-01	M18878_1	NT	Human diabifidin 27 gene, exons 10 and 11, and L1 and Alu repeats
8845	22480		0.58	3.8E-01	11485620	NT	Porphyra purpurea mitochondrial, complete genome
100688	22714	369332	0.77	3.8E-01	D86722_1	NT	Nicotiana tabacum mRNA for TATA binding protein (TBP), complete cds
101722	23410		1.98	3.8E-01	AV695974	EST_HUMAN	AV695974 GKC Homo sapiens cDNA clone GKCBQC11 5'
11753	24344	37674	1.47	3.8E-01	AV702623	EST_HUMAN	AV702623 ADB Homo sapiens cDNA clone ADDBDBE08 6'
119448	25285		3.37	3.8E-01	AF304354_1	NT	Homo sapiens proteoglycan 3 (PRC3) gene, complete cds
120688	24881		2.08	3.8E-01	Q61670	SWISSPROT	HOMEBOX PROTEIN HLX1
125589	24891		1.44	3.8E-01	11493335	NT	Homo sapiens hypothetical protein FLJ10583 (FLJ10583), mRNA
156	12871		8.33	3.8E-01	7018498	NT	Homo sapiens protein kinase PKNbeta (pkinbeta), mRNA
1863	14601		1.03	3.8E-01	AE003870_1	NT	Xylella fastidiosa, section 16 of 229 of the complete genome
2460	15178	27918	1.29	3.8E-01	U41846_1	NT	Ceanorhabditis briggsae acetylecholinesterase (ace-1) gene, complete cds
2576	15280	28027	1.62	3.8E-01	AF214117_1	NT	Arabidopsis thaliana putative c-myb-like transcription factor (MYB3R-3) mRNA, complete cds
2638	15691	28062	3.98	3.8E-01	66780002	NT	Mus musculus scilis carrier family 1 member 6 (Slc6a6), mRNA
3033	15769		1.14	3.8E-01	AJ251057_1	NT	Human immunodeficiency virus type 1 complete genome (Isolate 88SE-MP1213)
3043	15809	28456	1.39	3.8E-01	AF043383_1	NT	Plautelectes amhericenus aminopeptidase N (ampn) gene, partial cds
3477	16233	28887	7.98	3.8E-01	AL161618_2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 30
5527	16283		0.79	3.8E-01	AI807219_1	EST_HUMAN	Wf38p12_x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2257855 3'
5541	16283		1.22	3.8E-01	AI807218_1	EST_HUMAN	Wf38p12_x1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2257856 3'
3739	16492	29127	1.16	3.8E-01	BE1640B0_1	EST_HUMAN	PM04HT0339-20040-010-G01 HT0339 Homo sapiens cDNA
3897	16847	29287	0.97	3.8E-01	8754095	NT	Mus musculus general transcription factor II (Gtf2), mRNA
4043	16788	29416	0.74	3.8E-01	AJ271381_2	NT	Takifugu rubripes wrf2 (partial), transk1, crt and transk2 (partial) genes
6522	16820	31221	1.42	3.8E-01	Q04888	SWISSPROT	TRANSCRIPTION FACTOR SOX-10
6247	19021		0.74	3.8E-01	S46825_1	NT	Prion protein [mink, Genomic, 2446 nt]
6528	19294	32298	5.5	3.8E-01	BE072399_1	EST_HUMAN	QV3-B10537-271289-048-302 BT0537 Homo sapiens cDNA
6882	18579	32874	4.53	3.8E-01	AJ374601_1	EST_HUMAN	tc54f11_x1 Soares_tcl1_fusus_Nb2H-F8_8w Homo sapiens cDNA clone IMAGE:2047817 3' similar to contains Alu repetitive element;
6840	18502	32527	1.25	3.8E-01	AL161533_2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 25
7418	20093		4.42	3.8E-01	X671597_1	NT	M.musculus gene for klf11/klf11-binding protein
8198	20890	34029	0.86	3.8E-01	M81385_1	NT	M.musculus gene for klf11/klf11-binding protein (LRF-1)mRNA, complete cds
8455	21147	34229	2.04	3.8E-01	AB046835_1	NT	Mouse liver receptor homologous protein (LRH-1)mRNA, partial cds
8523	21215	34368	1.02	3.8E-01	11441284	NT	Homo sapiens FOS-like antigen-1 (FOXA1), mRNA
8716	21408	34651	1.28	3.8E-01	AL163279_2	NT	Homo sapiens chromosome 21 segment HS21C079
9461	22011		3.95	3.8E-01	TB5413_1	EST_HUMAN	ye33b03_x1 Soares fetal liver spleen cDNA clone IMAGE:120539 5' similar to contains Alu repetitive element; contains PTR5 repetitive element;

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Table 4  
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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar BLAST E Value	Top Hit No.	Top Hit Database Source	Top Hit Descriptor
10395	23396			1.67	3.8E-01	AY765814.1	EST_HUMAN
11521	24121			3.18	3.8E-01	BE710219.1	EST_HUMAN
11693	24288	37610		2.27	3.8E-01	R42550.1	EST_HUMAN
11693	24288	37611		2.27	3.8E-01	R42550.1	EST_HUMAN
12149	24636			4.76	3.8E-01	AE001124.1	NT
12270	25316			2.08	3.8E-01	U94788.1	NT
12384	24779			3.39	3.8E-01	BE829258.1	EST_HUMAN
12723	24994			1.54	3.8E-01	U78631.1	NT
12771	25201			1.74	3.8E-01	AF281483.1	NT
12788	25640	30986		1.51	3.8E-01	AF184972.1	NT
2488	15203	27844		12.24	3.7E-01	AB037831.1	NT
3453	16209	28860		9.84	3.7E-01	AF056336.1	NT
4284	16945	28572		7.39	3.7E-01	AI218707.1	EST_HUMAN
4285	17025	28651		1.3	3.7E-01	AW878037.1	EST_HUMAN
4357	17085	28730		2.65	3.7E-01	AE002468.1	NT
5678	18470	31398		1.19	3.7E-01	AF135187.1	NT
5860	18847	31688		0.9	3.7E-01	AI183278.2	NT
6417	19185	32183		0.88	3.7E-01	M10806.1	NT
6436	19204			0.72	3.7E-01	L10363.1	NT
7043	19734	32794		3.23	3.7E-01	11626843	NT
7695	20349	33463		0.8	3.7E-01	768802.1	EST_HUMAN
7719	20383	33497		0.56	3.7E-01	AW511328.1	EST_HUMAN
8227	20921	34059		2.07	3.7E-01	11436739	NT
8227	20921	34060		2.07	3.7E-01	11436739	NT
8268	20967	34096		0.65	3.7E-01	AA902812.1	EST_HUMAN
9101	21789			1.31	3.7E-01	AJ271386.1	NT
10089	22717			0.6	3.7E-01	K00861.1	NT
10110	22758	35970		4.12	3.7E-01	AI330411.1	EST_HUMAN
10184	23448			1.98	3.7E-01	X05958.1	NT
10957	23633	36882		2.81	3.7E-01	AJ297357.1	NT
10957	23633	36883		2.81	3.7E-01	AJ297357.1	NT
11443	23210	36441		2.75	3.7E-01	X04422.1	NT
11676	24271	37693		1.43	3.7E-01	D78248.1	EST_HUMAN

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Table 4  
Single Exon Probes Expressed In Brain

Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
11771	24362		2.87	3.7E-01	6877678	NT	<i>Mus musculus</i> retinoblastoma 1 (Rb1), mRNA
11869	24943		2.11	3.7E-01	J04982.1	NT	Human heart/skeletal muscle ATP/ADP translocator (ANT1) gene, complete cds
12033	24558		3.08	3.7E-01	AJ243325.1	NT	Chlamydomonas psittaci partial cmp1 gene for outer membrane protein 1
12488	24847		1.9	3.7E-01	AL121154.1	EST_HUMAN	DKFZp62K075_11762 (synonym: hmel2) Homo sapiens cDNA clone DKFZp762K075_5
12548	24886		3.08	3.7E-01	Y18000.1	NT	Homo sapiens NF2 gene
254	13082	25701	2.17	3.6E-01	AJ009809.1	NT	Brassica napus mRNA for MAP4K alpha2 protein
975	13740		8.22	3.6E-01	U89241.1	NT	Human mbp gene, partial cds
1281	14040	28713	3.83	3.6E-01	780255.1	EST_HUMAN	ye03ad05_r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:24443_5'
1281	14040	28714	3.93	3.6E-01	780256.1	EST_HUMAN	ye03ad05_r1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:24443_5'
18019	14648	27356	6.73	3.6E-01	AW590184.1	EST_HUMAN	hg33tf02.x1 NCI_CGAP GC8 Homo sapiens cDNA clone IMAGE:2847419_3'
18019	14848	27357	6.73	3.6E-01	AW590184.1	EST_HUMAN	hg33tf02.x1 NCI_CGAP GC8 Homo sapiens cDNA clone IMAGE:2847419_3'
1844	14679	27393	5.7	3.6E-01	AF216207.1	NT	<i>Mus musculus</i> ribosomal protein S10 (Rps10) gene, complete cds
2047	14780		1.39	3.6E-01	AF056827.1	NT	Rattus norvegicus repeat element associated with the Ratsnfrf gene
2267	14883		1.05	3.6E-01	AB002321.1	NT	Human mRNA for KIAA0323 gene, partial cds
2389	15110		2.69	3.6E-01	X76725.1	NT	P irregularis (Psp04) gene for actin
2479	15187	27938	1.23	3.6E-01	LD5436.1	NT	Rattus norvegicus synaptic vesicle protein (SV2) mRNA, complete cds
2479	15197	27937	1.23	3.6E-01	LD5435.1	NT	Rattus norvegicus synaptic vesicle protein (SV2) mRNA, complete cds
2491	15208	27950	1.43	3.6E-01	AW812033.1	EST_HUMAN	RC5-ST071-181098-01-907 ST0711 Homo sapiens cDNA
							PROTEIN-L-SOASPARTATE O-METHYLTRANSFERASE (PROTEIN-BETA-ASPARTATE METHYLTRANSFERASE) (PMT) (PROTEIN CARBOXYL METHYLTRANSFERASE) (L-SOASPARTATE ISOASPARTYL PROTEIN CARBOXYL METHYLTRANSFERASE)
2836	15348	28090	1.44	3.6E-01	P24206	SWISSPROT	Drosophila melanogaster sugar transporter 3 (su3) mRNA, complete cds
2800	17884		7.16	3.6E-01	AF109485.1	NT	H.sapiens serotonin transporter gene, exons 9 and 10
3462	16218	28871	2.16	3.6E-01	X79758.1	NT	H.sapiens serotonin transporter gene, exons 9 and 10
3462	16218	28872	2.16	3.6E-01	X79758.1	NT	H.sapiens serotonin transporter gene, exons 9 and 10
4375	17112	28745	'3	3.6E-01	BE707883.1	EST_HUMAN	RC1+HT0845_150880-014-b12 HT0845 Homo sapiens cDNA
4948	17676	30285	2.38	3.6E-01	AW338983.1	EST_HUMAN	hs02-g04.x1 NCI_CGAP Lu24 Homo sapiens cDNA clone IMAGE:2872666_3'
62288	18103	30762	0.82	3.6E-01	AJ006585.1	NT	Homo sapiens lipase intran 6
5995	18778	31738	0.85	3.6E-01	P16431	SWISSPROT	FORMATE HYDROGENLYASE SUBUNIT 6 PRECURSOR (FH1 SUBUNIT 5) (HYDROGENASE 3 COMPONENT E)
6388	19155	32154	1.74	3.6E-01	Y10186.1	NT	Homo sapiens PHEX gene
7048	19739		32	3.6E-01	RFx090.1	EST_HUMAN	y774sd6_r1 Soares fetal liver spleen 1NFSL Homo sapiens cDNA clone IMAGE:275987_5'
7183	19869	32843	1.9	3.6E-01	AW027174.1	EST_HUMAN	wf72c10.x1 Soares_thymus_NHFTt Homo sapiens cDNA clone IMAGE:2513010_3' similar to TR:015117
8123	20817	33953	0.58	3.6E-01	P38167	SWISSPROT	O15117 FYN BINDING PROTEIN [1]; SCO-SPONDIN

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Probe Seq ID No:	Exon Seq ID No:	ORF Seq ID No:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
8177 20871	34005		11.45	3.6E-01	AL161583.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 79
8890 21591	34731		2.74	3.6E-01	4504958	NT	Homo sapiens lysosomal-associated membrane protein 2 (LAMP2), transcript variant LAMP2A, mRNA
8900 21691	34732		2.74	3.6E-01	4504958	NT	[Homo sapiens lysosomal-associated membrane protein 2 (LAMP2), transcript variant LAMP2A, mRNA]
9091 21730	34944		1.17	3.6E-01	AL163204.2	NT	[Homo sapiens chromosome 21 segment HS2TC004
9259 21986	35139		1.04	3.6E-01	X17680.1	NT	D. melanogaster striped gene, exons 3, 4, 5 & 6
9259 21986	35140		1.04	3.6E-01	X17680.1	NT	D. melanogaster striped gene, exons 3, 4, 5 & 6
9269 21944			0.57	3.6E-01	X02525.1	NT	C. perfringens plc gene for phospholipase C upstream region containing bent DNA fragment
9763 22414	35621		14.67	3.6E-01	Q63184	SWISSPROT	PROBABLE PEPTIDE ABC TRANSPORTER ATP-BINDING PROTEIN Y413
9893 22543	35735		0.61	3.6E-01	AW752801.1	EST_HUMAN	MR2_CTD0222-21 1089-002-b10 CT0222 Homo sapiens cDNA
9893 22543	35736		0.51	3.6E-01	AW752801.1	EST_HUMAN	MR2_CTD0222-21 1089-002-b10 CT0222 Homo sapiens cDNA
10864 23544	36791		3.31	3.6E-01	BE02390.1	EST_HUMAN	60167641 AF1 NIH_MGC_21 Homo sapiens cDNA clone IMAGE:3855897 5'
11062 23722	36983		4.12	3.6E-01	AB004283.1	NT	Arabidopsis thaliana mRNA for SlgB, complete cds
11421 23188	36419		3.4	3.6E-01	AE0000856.1	NT	Methanobacterium thermophilum totophum from bases 702375 to 714311 (section 62 of 148) of the complete genome
11603 26416			1.83	3.6E-01	Y18210.1	NT	Homo sapiens Hifb5 gene for hair keratin, exons 1 to 9
11978 24522			1.4	3.6E-01	DB0001.1	NT	Synecocystis sp. PCC6803 complete genome, 3/27, 27160-402289
11987 24523			3.89	3.6E-01	AE0000335.1	NT	Escherichia coli K-12 MG1655 section 225 of 400 of the complete genome
12135 24624			4	3.6E-01	U66888.1	NT	Mus musculus Emr1 mRNA, complete cds
12483 24850			2.12	3.6E-01	11432598	NT	Homo sapiens myeloid lymphoid or mixed-lineage leukaemia (Drosophila homolog); translocated to, 10 (AF10), mRNA
12748 25383			2.23	3.6E-01	AW190229.1	EST_HUMAN	x60911.X1 NC_ OGAP_Par1 Homo sapiens cDNA clone IMAGE:2579116 3' similar to gatk005538 TUBULIN ALPHA-1 CHAIN (HUMAN)
204 13017	25657		2.05	3.5E-01	68778533	NT	Mus musculus mannose receptor, C type 2 (Mrc2), mRNA
708 13482	28131		1.69	3.5E-01	7706138	NT	Homo sapiens GAP-like protein (LOC51306), mRNA
708 13482	28132		1.59	3.5E-01	7706138	NT	Homo sapiens GAP-like protein (LOC51306), mRNA
782 13535	28194		4.25	3.5E-01	BF128786.1	EST_HUMAN	6018110601R1 NIH_MGC_48 Homo sapiens cDNA clone IMAGE:4053951 3'
1815 14362	27053		1.1	3.5E-01	BF310588.1	EST_HUMAN	6018046531F2 NIH_MGC_18 Homo sapiens cDNA clone IMAGE:4124244 5'
1636 14362	27068		1.96	3.5E-01	U35776.1	NT	Rattus norvegicus ADP-ribosylation factor-directed GTPase activating protein mRNA, complete cds
2281 15006	27747		1.35	3.5E-01	P06798	SWISSPROT	HOMEBOX PROTEIN HOXA4 (HOX-1.4) (MH-3)
2812 15600	28066		1.76	3.5E-01	AA22252.1	EST_HUMAN	270809.s1 Strategene NT2 neuronal precursor 937230 Homo sapiens cDNA clone IMAGE:6509872 3'

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
3785 16537			0.86	3.6E-01	AA642138.1	EST_HUMAN	nr60d03_s1 NCI_CGAP_Lym3 Homo sapiens cDNA clone IMAGE:11723573'
4281 16972	28598	1.67	3.5E-01	AF071253.1	NT	Danio rerio homeobox protein (hoxb5b) gene, complete cds	
4443 17179	29895	0.94	3.6E-01	BE146885.1	EST_HUMAN	RC5-HT0218-181088-011-902 HT0218 Homo sapiens cDNA	
4627 17362	29895	1.02	3.6E-01	Y18477.1	NT	Mus musculus Axotrophin gene 5' flanking region	
4880 17601	30230	4.58	3.5E-01	MA18348.1	NT	Rat leukocyte common antigen (L-C4) gene, exons 1 through 5	
5251 18057	30685	0.76	3.6E-01	Q86687	SWISSPROT	EARLY EA DNA-BINDING PROTEIN	
5251 18057	30686	0.76	3.6E-01	Q86687	SWISSPROT	EARLY EA DNA-BINDING PROTEIN	
5462 18281	31152	1.13	3.6E-01	DA2045.1	NT	Human mRNA for KIAA0038 gene, complete cds	
6143 18821		0.98	3.6E-01	AW883198.1	EST_HUMAN	PM4-SN012-030400-001-011 SN0012 Homo sapiens cDNA mRNA, complete cds	
6314 19085	32070	0.6	3.6E-01	AA431833.1	EST_HUMAN	ZW78f03_11 Scores_tests_NHT Homo sapiens cDNA clone IMAGE:782429 5' similar to TR-G10868935	
6359 19129	32124	0.68	3.6E-01	U37150.1	NT	Bos taurus peptide methionine sulfide reductase (msrA) mRNA, complete cds	
6566 18831	32338	1.08	3.6E-01	O24287	SWISSPROT	GLUCOSE-6-PHOSPHATE 1-DEHYDROGENASE, CHLOROPLAST PRECURSOR (CPD)	
6956 18438		4.24	3.6E-01	X985805.1	NT	S. scrofa mRNA for CD31 protein (PECAM-1)	
7441 20118	33207	0.55	3.5E-01	P47281	SWISSPROT	HISTIDYL-TRNA SYNTHETASE (HISTIDINE-TRNA LIGASE) (HISRS)	
7441 20118	33208	0.66	3.6E-01	P47281	SWISSPROT	HISTIDYL-TRNA SYNTHETASE (HISTIDINE-TRNA LIGASE) (HISRS)	
7970 20685		2.19	3.6E-01	11448302	NT	Homo sapiens tumor protein p53-binding protein, 2 (TP53BP2), mRNA	
7973 20688	33790	0.71	3.6E-01	BF358871.1	EST_HUMAN	RC4-E10/024-260800-014-d07 ET0024 Homo sapiens cDNA	
8368 21059		0.63	3.6E-01	AF051661.1	NT	Rattus norvegicus Na-K-Cl cotransporter (Nkcc1) mRNA, complete cds	
8825 21517	34662	1.17	3.6E-01	4507610	NT	Homo sapiens tyrosine kinase non-receptor 1 (TNK1), mRNA	
9636 22288	35481	1.52	3.6E-01	Q102294	SWISSPROT	VOLTAGE-DEPENDENT N-TYPE CALCIUM CHANNEL ALPHA-1B SUBUNIT (CALCIUM CHANNEL, L TYPE, ALPHA-1 POLYPEPTIDE ISOFORM 6) (BRAIN CALCIUM CHANNEL III) (BII)	
9786 22437	35644	5.94	3.6E-01	Z26825.1	NT	X. laevis gene for albumin including HIF-1 enhancer	
9887 22517	35713	0.98	3.6E-01	BE174794.1	EST_HUMAN	QV2-HT0577-090400-122-007 HT0577 Homo sapiens cDNA	
10635 23327	36564	2.78	3.6E-01	X61084.1	NT	C. elegans rho50kDa gene for opsin protein	
10948 23625	36875	2.39	3.6E-01	AJ243178.1	NT	Gallus gallus SPARC gene for osteonectin, promoter and exon 1	
10948 23625	36876	2.39	3.6E-01	AJ243178.1	NT	Gallus gallus SPARC gene for osteonectin, promoter and exon 1	
11505 24106	37419	1.34	3.5E-01	U07000.1	NT	Human breakpoint cluster region (BCR) gene, complete cds	
11695 24184	37499	1.64	3.5E-01	N77587.1	EST_HUMAN	y80h12_r1 Scores_multiple_scenes_2NhlhSP_Homo sapiens cDNA clone IMAGE:2803765'	
11695 24216		1.71	3.6E-01	M82885.1	NT	Drosophila melanogaster dual bar protein (BarH2) gene, exon 1	
11694 24279	37601	1.51	3.5E-01	L05145.1	NT	Human glucokinase (GCK) gene, repeat polymorphism	
11776 24367		1.36	3.5E-01	A1084773.1	EST_HUMAN	HA0542 Human fetal liver cDNA library Homo sapiens cDNA	
12063 24578		1.47	3.6E-01	X64585.1	NT	Streptomyces etiA1 gene for F(0)F(1)ATP synthase alpha-subunit	
12214 24876		2.32	3.6E-01	AE001774.1	NT	Thermotoga maritima section 86 of 136 of the complete genome	

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
12402	24787			1.4	3.5E-01 AE001697.1	NT	Thermitoga maritima section 3 of 138 of the complete genome
12783	25269	30723	3.33	3.5E-01 H80814.1	EST_HUMAN	ye64f11_r1 Soenes retina N2b4HR Homo sapiens cDNA clone IMAGE:218697 5'	
12783	25269	30724	3.33	3.5E-01 H80814.1	EST_HUMAN	ye64f11_r1 Soenes retina N2b4HR Homo sapiens cDNA clone IMAGE:218697 5'	
691	13468			1.85	3.4E-01 AJ242258.1	NT	Homo sapiens partial N-myc (exon 3), HPV45 L2, HPV45 E6, HPV45 E7 and HPV45 E1 genes isolated from IC4 cervical carcinoma cell line
955	13720	26898	7.61	3.4E-01 Y097862	NT	Pseudomonas fluorescens ccrR, ccrS genes, oriC222 and partial lacA gene	
1363	14052	26725	1.72	3.4E-01 Y00554.1	NT	Azotobacter vinelandii nifA gene for NifA protein (positive regulatory element)	
2400	15121	27858	2.92	3.4E-01 D80909.1	NT	Synechocystis sp. PCC6803 complete genome, 11/27, 1311235-1430418	
39001	15707	28416	0.85	3.4E-01 AL163210.2	NT	Homo sapiens chromosome 21 segment HS21C210	
39001	15707	28416	0.85	3.4E-01 AL163210.2	NT	Homo sapiens chromosome 21 segment HS21C210	
31448	15910	26555	1.08	3.4E-01 D80909.1	NT	Synechocystis sp. PCC6803 complete genome, 11/27, 1311235-1430418	
3159	15922	28588	6.23	3.4E-01 U83905.1	NT	Canis familiaris rod photoreceptor cGMP-gated channel alpha-subunit (CNGC1) mRNA, complete cds	
33238	16098	28749	0.9	3.4E-01 AF034882.1	NT	Homo sapiens pulmonary surfactant protein D, promoter region and exon 1	
35222	16278	28933	3.48	3.4E-01 AF106835.1	NT	Methylyavorus sp. strain SS1 putative GpE (gpe), DnaK (dnak), and putative DnaJ (dnaj) genes, complete cds	
3770	16522			1.69	3.4E-01 BF449010.1	EST_HUMAN	789401_r1 NCI_OGAP_Ov18 Homo sapiens cDNA clone IMAGE:3572232 3' similar to TRQ8UJ16 Q8WU16 DJ18C9.1
40229	16774			2.38	3.4E-01 AA5B498.1	EST_HUMAN	nt011b10_s1 NCI_OGAP_Phe1 Homo sapiens cDNA clone IMAGE:7100347 3'
2480	17198	28923	0.82	3.4E-01 AF1683341.1	NT	Homo sapiens integrin alpha 8 (ITGA8) gene, exons 12 through 23	
4599	17334	28933	1.54	3.4E-01 BE080912.1	EST_HUMAN	MR4-BT0403-230200-202-01 BT0403 Homo sapiens cDNA qj8605_x1 NCI_OGAP_Kid3 Homo sapiens cDNA clone IMAGE:1887208 3' similar to contains Alu repetitive element	
4898	17825		3.23	3.4E-01 AI240973.1	EST_HUMAN		
6143	17862			0.98	3.4E-01 U78740.1	NT	Homo sapiens serotonin transporter (SERT) gene, promoter region, exons 1B and 2, and partial cds
5599	18394	31304	2.62	3.4E-01 AL161594.2	NT	Arachidopsis thaliana DNA chromosome 4, contig fragment No. 90	
6721	18613		6.09	3.4E-01 AA085313.1	EST_HUMAN	zn12a11_s1 Strategene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:547221 3'	
5917	18702			1.89	3.4E-01 LD2871.1	NT	Echovirus 22_1AB_1C_1D_2A_2B_2C_3A_3B_3C_3D proteins RNA, complete mature peptides and cds
6940	18722	31681	0.89	3.4E-01 BE748912.1	EST_HUMAN	601571811T1 NIH_MGC_55 Homo sapiens cDNA clone IMAGE:3838828 3'	
6917	18798	31769	2.43	3.4E-01 AW204905.1	EST_HUMAN	UH-B1-8el-8-12-0-ULs1 NCI CGAP_Sub3 Homo sapiens cDNA clone IMAGE:2718582 3'	
6141	18919	31889	1.81	3.4E-01 AL120544.1	EST_HUMAN	DKFZp761A249_r761 (synonym: ham2) Homo sapiens cDNA clone DKFZp761A249 5'	
6844	19466		1.56	3.4E-01 N85226.1	EST_HUMAN	zb53e12_s1 Soenes_fetal_lung_Nbh119W Homo sapiens cDNA clone IMAGE:307342 3'	

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Table 4  
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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
6848	18548	32578	1.02	3.4E-01	AI468082.1	EST_HUMAN	Im63g05_x1 NCI_CGAP_Bm25 Homo sapiens cDNA clone IMAGE:2162840 3' similar to gb:S37431
68589	19441	32458	0.59	3.4E-01	BF678702.1	EST_HUMAN	LAMININ RECEPTOR (HUMAN); 602085283FT NIH MCG_83 Homo sapiens cDNA clone IMAGE:2248365 5'
78906	20501		0.49	3.4E-01	AE000493.1	NT	Escherichia coli K-12 MG1655 section 383 of 400 of the complete genome
81395	20829	33894	0.6	3.4E-01	Y14930.1	NT	Homo sapiens TCRAV28 gene, allele A4, partial
81488	20882		0.47	3.4E-01	BF448010.1	EST_HUMAN	7194a01_x1 NCI_CGAP_Ov18 Homo sapiens cDNA clone IMAGE:3572232 3' similar to TRQ9UJ16
83886	21079		1.51	3.4E-01	AA337063.1	EST_HUMAN	EST141765 Endometrial tumor Homo sapiens cDNA 5' end
84601	21153	34286	0.72	3.4E-01	LD4680.1	NT	Cricetulus griseus cholesterol 7-alpha-hydroxylase gene, complete cds
87571	21443	34580	1.7	3.4E-01	8833624	NT	Bovine enterovirus strain K2577, complete genome
91112	21800	34964	4.42	3.4E-01	P26013	SWISSPROT	INTEGRIN BETA-8 PRECURSOR
91112	21800	34965	4.42	3.4E-01	P26013	SWISSPROT	INTEGRIN BETA-8 PRECURSOR
93221	21988		0.61	3.4E-01	AB017610.1	NT	Ephydite flavifolia mRNA for PLC-gammaS, complete cds
93446	20417	33535	4.67	3.4E-01	U19482.1	NT	Saccharomyces cerevisiae Maf1p (MAF1) gene, complete cds
93446	20417	33537	4.87	3.4E-01	U19482.1	NT	Saccharomyces cerevisiae Maf1p (MAF1) gene, complete cds
83397	220159	35229	0.5	3.4E-01	AF183857.1	NT	Dictyostelium discoideum putative Chk/F receptor CMFR1 mRNA, complete cds
65995	22248	35433	1.01	3.4E-01	U687783.1	NT	Glycine max putative transcription factor SCOF-1 (soof-1) mRNA, complete cds
87789	22440	35648	1.83	3.4E-01	AJ225034.1	NT	Homo sapiens FAA gene, exon 16, 17 and 18
103376	23022		0.62	3.4E-01	AE004088.1	NT	Vibrio cholerae chromosome I, section 4 of 261 of the complete chromosome
10940	23620		4.72	3.4E-01	AE000881.1	NT	Methanobacterium thermophilicum from busses 1018444 to 1028212 (section 87 of 148) of the complete genome
10984	23659	36912	2.6	3.4E-01	P08925	SWISSPROT	PROBABLE E4 PROTEIN
11032	23703	36971	2.17	3.4E-01	AF045881.1	NT	Rutilus arvensis cytochrome b (cytb) gene, mitochondrial gene encoding mitochondrial protein, partial cds
11253	23915	37207	1.81	3.4E-01	M28858.1	NT	Human von Willebrand factor gene, exons 36 and 37
11253	23916	37208	1.81	3.4E-01	M28858.1	NT	Human von Willebrand factor gene, exons 36 and 37
11463	24084	37386	1.88	3.4E-01	AB035507.1	NT	Rattus norvegicus mRNA for e-glicerin MUC18, complete cds
11513	24113	37423	3.85	3.4E-01	AL161616.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 27
11798	24378	37708	1.72	3.4E-01	BF061948.1	EST_HUMAN	7169d12_x1 NCI_CGAP_GC8 Homo sapiens cDNA clone IMAGE:3480846 3'
11861	24445	37786	1.98	3.4E-01	U07000.1	NT	Human breakpoint cluster region (BCR) gene, complete cds
11881	24455		1.85	3.4E-01	U83604.1	NT	Citrus variegation virus putative replicase gene, partial cds
12197	24668		11.43	3.4E-01	L26839.1	NT	Human antigenigen mRNA, complete cds
12224	25192		1.61	3.4E-01	BE218652.1	EST_HUMAN	hu42d08_x1 NCI_CGAP_LW24 Homo sapiens cDNA gene IMAGE:3170127 3' similar to controls PTR5.3 PTR5 repetitive element;

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
12280	25282		2.28	3.4E-01	8893361	NT	Beta vulgaris mitochondrial, complete genome
12391	24781	31036	2.2	3.4E-01	AJ287131.1	NT	Mus musculus SII, MAP_17, CYP_4, SCL & CYP_b genes
12888	24974		1.82	3.4E-01	AF018413.1	NT	Homo sapiens HLA class III region containing tenascin X (tenascin-X) gene, partial cds; cytochrome P450 21-hydroxylase (CYP21B), complement component C4 (C4B) G11, hemicase (SK12W), RD, complement factor B (Bf), and complement component C2 (C2) genes. >
13	12840	25453	10.77	3.3E-01	X07890.1	NT	Rhizobium leguminosarum sym plasmid pRL5.11 nodX gene
103	12840	25453	4.4	3.3E-01	X07890.1	NT	Rhizobium leguminosarum sym plasmid pRL5.11 nodX gene
435	13221	25887	0.9	3.3E-01	AL101545.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 45
618	13397	26032	2.01	3.3E-01	7682485	NT	Homo sapiens KIAA1100 protein (KIAA1100), mRNA
1178	13631	26597	2.85	3.3E-01	Q12448	SWISSPROT	PROLINE-RICH PROTEIN LAS17
1284	14034	28705	3.76	3.3E-01	BF568880.1	EST_HUMAN	60218401671 NIH_MGC_42 Homo sapiens cDNA clone IMAGE:4300251 3'
1336	14085	28780	1.2	3.3E-01	U43626.1	NT	Human chromosome 15q11-q13 putative DNA replication origin in the g-aminobutyric acid receptor b3 and a5 gene cluster
1601	14347	27036	1.47	3.3E-01	6763665	NT	Mus musculus diisogelin 3 (Dtnn5), mRNA
1731	14473		1.02	3.3E-01	AA332734.1	EST_HUMAN	EST38722 Embryo, 8 week Homo sapiens cDNA 5' end
2022	14757		1.01	3.3E-01	AF081148.1	NT	Methylococcus capsulatus strain Bath outer membrane protein MapB (mapB) gene, complete cds
2404	16125		4.62	3.3E-01	480784	NT	Homo sapiens uridine monophosphate synthetase (uridylate phosphorylase) transferase and carboxypeptidase (carboxypeptidase) (UMPS), mRNA
2849	16715	28368	1.87	3.3E-01	AJ251805.1	NT	Bacteriophage phi-Yac3-12 complete genome
3051	15817	28462	1.48	3.3E-01	AJ007822.2	NT	Streptomyces erythraea mithramycin biosynthetic genes
3488	16243	28699	1.07	3.3E-01	AB012822.1	NT	Homo sapiens MTA1-L1 gene, complete cds
3789	16541	29178	2.1	3.3E-01	O84645	SWISSPROT	EXODEOXYRIBONUCLEASE V BETA CHAIN
3789	16551	29183	0.97	3.3E-01	P22602	SWISSPROT	GENOME POLYPROT [CONTAINS: N-TERMINAL PROTEIN (P1); HELPER COMPONENT PROTEINASE (HC-PRO); PROTEIN P3]
3932	16882	28323	1.03	3.3E-01	4757739	NT	Homo sapiens A kinase (PRKA) inhibitor protein 5 (AKAP5), mRNA
3947	16897	28338	1.47	3.3E-01	AL161488.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 10
3983	16731	28385	1.79	3.3E-01	AF200446.1	NT	Hypoxylon fragiforme chitin synthase gene, partial cds
4334	17073		1.8	3.3E-01	D31682.1	NT	Reitius nonnegicetus DNA for regucalcin, partial cds
4641	17376		1.23	3.3E-01	AI538114.1	EST_HUMAN	tp78b12L1 NCI CGAP_113 Homo sapiens cDNA clone IMAGE:2205407 3' similar to g:X57522 ANTIGEN PEPTIDE TRANSPORTER 1 (HUMAN)
4798	17517	30139	1.22	3.3E-01	D64003.1	NT	Synecchocystis sp. PCC6803 complete genome, 22/27, 27/55/03-28/68/78
5146	17885		0.98	3.3E-01	AW837982.1	EST_HUMAN	QVO-D70047-170200-123-H08 D70047 Homo sapiens cDNA
5241	18047	30675	2.61	3.3E-01	289819.1	NT	R.noventicis mRNA for 3'UTR of ubiquitin-like protein
5241	18047	30678	2.61	3.3E-01	289819.1	NT	R.noventicis mRNA for 3'UTR of ubiquitin-like protein

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
5700	18494	31417	0.74	3.3E-01	BF213873.1	EST_HUMAN	601848080F_NIH_MGC_55 Homo sapiens cDNA clone IMAGE-4078823 5'
66936	18843	31682	1.9	3.3E-01	BE618650.1	EST_HUMAN	601472768T_NIH_MGC_88 Homo sapiens cDNA clone IMAGE-3875753 3'
66938	18843	31683	1.9	3.3E-01	BE618650.1	EST_HUMAN	601472768T_NIH_MGC_88 Homo sapiens cDNA clone IMAGE-3876753 3'
5647	18729	31688	1.18	3.3E-01	P05691	SWISSPROT	CIRCUMSPOROZOITE PROTEIN (CS)
66936	19012	32851	0.71	3.3E-01	AB034233.1	NT	Flexibacter literalis gyB gene for DNA gyrase B subunit, partial cds
66935	18812	32852	0.71	3.3E-01	AB034233.1	NT	Flexibacter literalis gyB gene for DNA gyrase B subunit, partial cds
6789	19833	32560	4.82	3.3E-01	AI628131.1	EST_HUMAN	ty64h01.x1_NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE-2285809 3' similar to contains Alu repetitive element;contains element L1 repetitive element;
6789	19833	32561	4.82	3.3E-01	AI628131.1	EST_HUMAN	ty64h01.x1_NCI_CGAP_Kid11 Homo sapiens cDNA clone IMAGE-2285809 3' similar to contains Alu repetitive element;contains element L1 repetitive element;
7682	20346	33458	1.68	3.3E-01	N85148.1	EST_HUMAN	J2468F Human fetal heart, Lambda ZAP Express Homo sapiens cDNA clone J2498 5' similar to TEGT
8460	21152	34295	18.62	3.3E-01	BF883854.1	EST_HUMAN	602140372F_NIH_MGC_48 Homo sapiens cDNA clone IMAGE-4301800 5'
8859	21351	34497	0.48	3.3E-01	AU126115_NT2RP1	Homo sapiens cDNA clone NT2RP1/000130 5'	Homo sapiens cDNA clone NT2RP1/000130 5'
8859	21351	34498	0.48	3.3E-01	AU126115_NT2RP1	Homo sapiens cDNA clone NT2RP1/000130 5'	Homo sapiens cDNA clone NT2RP1/000130 5'
9012	21702	34852	0.81	3.3E-01	Q62225	SWISSPROT	MITOGEN-ACTIVATED PROTEIN KINASE KINASE KINASE 1 (MAP/ERK KINASE KINASE 1) (MEKK1) KINASE 1 (MEKK1)
9278	22032	35203	0.81	3.3E-01	BE828481.1	EST_HUMAN	CN3-ET0041-180500-187-d10 ET0041 Homo sapiens cDNA
9278	22032	35204	0.81	3.3E-01	BE828481.1	EST_HUMAN	CN3-ET0041-180500-187-d10 ET0041 Homo sapiens cDNA
9411	22073	35244	2.62	3.3E-01	N86860.1	EST_HUMAN	2a67h01.61 Seares, fetal lung, NbH1.16W Homo sapiens cDNA clone IMAGE-287649 3'
9452	22012	35174	2.77	3.3E-01	BF576745.1	EST_HUMAN	RC4-TR077-250800-011-d04 TN0077 Homo sapiens cDNA
8891	22541		2.27	3.3E-01	L41044.1	NT	Homo sapiens high-mobility group phosphoprotein (HMGI-C) gene, exons 1-3, complete cds
10622	23315	36554	3.13	3.3E-01	X63953.1	NT	D.mauritiana Adh gene
10622	23315	36555	3.13	3.3E-01	X63953.1	NT	D.mauritiana Adh gene
10951	23623		1.7	3.3E-01	BF528499.1	EST_HUMAN	60207682F1_NCI_O3AP_Bm84 Homo sapiens cDNA clone IMAGE-4213585 5'
11198	23861	37147	11.61	3.3E-01	BE216851.1	EST_HUMAN	hf61g02x1_NCI_CGAP_Lu24 Homo sapiens cDNA clone IMAGE-3176978 3'
11317	24008	37313	3.23	3.3E-01	P47953	SWISSPROT	GALECTIN-3 (GALACTOSE-SPECIFIC LECTIN 3) (MAC-2 ANTIGEN) (IGE-BINDING PROTEIN) (35 kD LECTIN) (CARBOHYDRATE BINDING PROTEIN 35) (CBP 35) (LECTIN L-29) (CBP30)
11179	24913		3.06	3.3E-01	AA808621.1	EST_HUMAN	ab71g02.51_NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE-1336850 3'
11174	12840	26453	1.87	3.3E-01	X07880.1	NT	Rhizobium leguminosarum sym plasmid pRL5J nodX gene
11197	24621	37286	1.71	3.3E-01	65983319	NT	Homo sapiens aldehyde oxidase 1 (AOX1), mRNA
12076	24987		3.34	3.3E-01	AP0000021	NT	Pseudomonas aeruginosa O75 genomic DNA, 287001-54400 nt. position (277)
444	13230		2.33	3.2E-01	AF018261.1	NT	Rattus norvegicus EH domain binding protein Epsin mRNA, complete cds

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
701	13476		1.43	3.2E-01	AL167561.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 61
1139	13884	26855	27.53	3.2E-01	AF047013.1	NT	Fusarium roseum virus 1 RNA2 putative RNA dependent RNA polymerase gene, complete cds
1259	14008	26877	1.38	3.2E-01	Z50202.1	NT	P. vulgaris arc5-1 gene
1389	14117	26782	5.42	3.2E-01	Q48624	SWISSPROT	LACTOSE PERMEASE (LACTOSE-PROTON SYMPORT) (LACTOSE TRANSPORT PROTEIN)
1767	14569	27210	1.28	3.2E-01	Z36041.1	NT	S. cerevisiae chromosome II reading frame ORF YBR172c
1777	14519	27222	4.7	3.2E-01	AW057194.1	EST_HUMAN	EST369224 MAGE sequences, MAGD Homo sapiens cDNA
1777	14519	27223	4.7	3.2E-01	AW057194.1	EST_HUMAN	EST369224 MAGE sequences, MAGD Homo sapiens cDNA
1835	14574	27283	1.23	3.2E-01	AL111655.1	NT	Babylis diherce straif T4 cDNA library under conditions of nitrogen deprivation
2157	14887	27621	2.52	3.2E-01	BF283817.1	EST_HUMAN	601868804/F NIH_MGC_17 Homo sapiens cDNA clone IMAGE-41115125'
2543	15257	28051	2.01	3.2E-01	7710079	NT	Mus musculus Pbx/motif 1 homeobox (Pbxmot1), mRNA
2713	15420	28159	1.08	3.2E-01	AF080568.1	NT	Homo sapiens promyelocytic leukaemia zinc finger protein (PLZF) gene, complete cds
3594	16347		0.77	3.2E-01	D10872.1	NT	Human h NAT allele 3-2 gene for arylamine N-acetyltransferase
4305	17044	28669	0.91	3.2E-01	4759185	NT	Homo sapiens synaptophysin (SYN) mRNA
4383	17101	29736	1.52	3.2E-01	M18818.1	NT	Rabbit beta-like globin gene cluster encoding the epsilon, gamma, delta (psuedogene) and beta globin polypeptides, complete cds
4484	17200	29826	1.21	3.2E-01	Q10288	SWISSPROT	HYPOTHETICAL 81.7 KD PROTEIN CT13G.04C IN CHROMOSOME 1 PRECURSOR
4688	17422		6.7	3.2E-01	BF9838617.1	EST_HUMAN	60120819721 NIH_MGC_81 Homo sapiens cDNA clone IMAGE-42465055'
4828	17557	30179	1.17	3.2E-01	Q57081	SWISSPROT	CYTADHERENCE HIGH MOLECULAR WEIGHT PROTEIN 3 (CYTADHERENCE ACCESSORY PROTEIN 3) (ACCESSORY ADHESIN PROTEIN 3) (P88)
4965	17680	30299	0.74	3.2E-01	BE782748.1	EST_HUMAN	6014655971 NIH_MGC_67 Homo sapiens cDNA clone IMAGE-38587895
5190	17898	30621	3.28	3.2E-01	BE73984.1	EST_HUMAN	CM04H10569-060303-269-410 HT0569 Homo sapiens cDNA
6868	18655	31568	1.07	3.2E-01	27221.1	NT	Giardia intestinalis Pyruvate-flavodoxin oxidoreductase and flanking genes
6211	18988	31983	0.9	3.2E-01	AF016494.1	NT	Fugu rubripes gamma-aminobutyric acid receptor beta subunit gene, partial cds; 55kd erythrocyte membrane protein (P55), synaptic vesicle-associated integral membrane protein (VAMP-1), procollagen C-proteinase enhancer protein (PCOLCE) genes, complete cds
6501	19268	32268	0.84	3.2E-01	AV718037.1	EST_HUMAN	AV718037 FHTA Homo sapiens cDNA clone FHT7A/BH101 5'
6834	19396		1.09	3.2E-01	AB002289.1	NT	Human mRNA for KIAA0361 gene, KIAA0361 protein
7755	20451	33575	0.51	3.2E-01	AJ277681.1	NT	Homo sapiens partial LM01 gene for LIM domain only 1 protein, exon 1
8072	20768	33955	1.48	3.2E-01	M60266.1	NT	Rat ISO-estradiol responsive factor gene, complete cds
8164	20858	33990	0.45	3.2E-01	AJ231001.1	NT	Rattus norvegicus repeat; map NOS-D12Wact1
8285	20959	34098	14.41	3.2E-01	X02508.1	NT	H.sapiens gene fragment for acetylcholine receptor (AChR) alpha subunit exons 8, 9 and 3' flanking region
8288	20982	34103	13.78	3.2E-01	BF311635.1	EST_HUMAN	6018871071 NIH_MGC_19 Homo sapiens cDNA clone IMAGE-41268335 6
8361	21054		1.38	3.2E-01	AL161574.2	NT	Arabidopsis thaliana DNA chromosome 4, contig fragment No. 70

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Probe SEQ ID NO:	Exon SEQ ID NO:	ORF SEQ ID NO:	Expression Signal	Most Similar (Top) Hit BLAST E Value	Top Hit Accession No.	Top Hit Database Source	Top Hit Descriptor
8898	21091	34228	1.24	3.2E-01	BF246771.1	EST_HUMAN	601855580F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:4075627 5
8898	21091	34227	1.24	3.2E-01	BF246771.1	EST_HUMAN	601855580F1 NIH_MGC_57 Homo sapiens cDNA clone IMAGE:4075627 5
8471	21193	34308	2.65	3.2E-01	AE002015.1	NT	Diphococcus radiodurans R1 section 152 of 228 of the complete chromosome 1
8571	21263	34401	0.84	3.2E-01	U510268.1	NT	Oryctolagus cuniculus Ig H-chain pseudogene, V-region (VH8-a2) gene, partial cds
8571	21263	34402	0.84	3.2E-01	U510268.1	NT	Oryctolagus cuniculus Ig H-chain pseudogene, V-region (VH8-a2) gene, partial cds
8895	21656	34807	0.51	3.2E-01	AL163204.2	NT	Homo sapiens chromosome 21 segment HS21C004
8876	21656	34808	2.18	3.2E-01	M86511.1	NT	Human monocyte/s antigen CD14 (CD14) mRNA, complete cds
9048	21737	34894	0.05	3.2E-01	AF041828.1	NT	Homo sapiens G-phosphofructo-2-kinase/fructose-2,6-bisphosphatase (PF2K) gene, exons 12 and 13
9048	21737	34895	0.05	3.2E-01	AF041828.1	NT	Homo sapiens G-phosphofructo-2-kinase/fructose-2,6-bisphosphatase (PF2K) gene, exons 12 and 13
8894	22544	35737	3.33	3.2E-01	U44914.1	NT	Bonellia burdettieri plasmid cp32-2, spc and spC genes, complete cds; and unknown genes
10099	22747	36982	0.45	3.2E-01	BE328230.1	EST_HUMAN	Inv805.1x NCI_CGAP_LU24 Homo sapiens cDNA clone IMAGE:3181569 3'
10210	22858		3.41	3.2E-01	AB011389.1	NT	Homo sapiens gene for AF-8, complete cds
10568	23261	36498	3.94	3.2E-01	T03813.1	EST_HUMAN	EST04702 Fetal brain, Stratagene Cat#338200) Homo sapiens cDNA clone HFBD221
12010	25317		3.91	3.2E-01	L07288.1	NT	Drosophila melanogaster laminin A (Lam-A) mRNA, complete cds
12392	25374		1.44	3.2E-01	BE888846.1	EST_HUMAN	60150782DF1 NIH_MGC_71 Homo sapiens cDNA clone IMAGE:3909532 6'
12524	24871		4.21	3.2E-01	Q83217	SWISSPROT	ELONGATION FACTOR TU (EF-TU)
12855	24855		2.07	3.2E-01	L38874.1	NT	Homo sapiens decoy/dystylose deaminase gene, complete cds
12712	25354	30606	1.76	3.2E-01	BE385778.1	EST_HUMAN	60127548DF1 NIH_MGC_20 Homo sapiens cDNA clone IMAGE:3616748 6'
2677	15388	28128	2.89	3.1E-01	R18051.1	EST_HUMAN	yeast06_r1 Soares fetal liver spleen INFLS Homo sapiens cDNA clone IMAGE:125051 5' similar to gb:M64241 QM PROTEIN (HUMAN);
2702	16532	28145	3.39	3.1E-01	7681971	NT	Homo sapiens KIAA0174 gene product (KIAA0174), mRNA
2702	16532	28146	3.39	3.1E-01	7681971	NT	Homo sapiens KIAA0174 gene product (KIAA0174), mRNA
2862	15630		1.29	3.1E-01	AW028036.1	EST_HUMAN	hik0h08_r1 Soares_NFL_T_GBC_S1 Homo sapiens cDNA clone IMAGE:2975391 3'
3170	15933		3.35	3.1E-01	AB029059.1	NT	Mus musculus gene for Ser/Thr kinase KKIAIMRE, exon 6
3887	16637	28278	0.8	3.1E-01	AJ251586.1	NT	Daucus carota mRNA for transcription factor E2F (E2F gene)
4908	17638	30250	0.73	3.1E-01	AE003984.1	NT	Xylella fastidiosa, section 130 of 229 of the complete genome
6380	18190	30882	9.73	3.1E-01	AF176111.1	NT	Homo sapiens hepatocyte nuclear factor-3 alpha (HNF3A) gene, exon 1
6513	18311	31212	0.73	3.1E-01	P44132	SWISSPROT	HYPOTHETICAL PROTEIN H1238
5514	18312	31213	0.67	3.1E-01	Z74883.1	NT	S.cerevisiae chromosome XV reading frame ORF YOL141w
5524	18322		0.88	3.1E-01	Y13278.1	NT	Mus musculus mRNA for polycystin
6685	18478	31393	2.11	3.1E-01	AF184122.1	NT	Homo sapiens filamin 2 (FLN2) gene, exons 10 through 22
6191	25087	31942	0.59	3.1E-01	R94322.1	EST_HUMAN	yq41fd_r1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:198387 5'